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(54) Title: SCREENING COMPOUNDS FOR THE ABILITY TO ALTER THE PRODUCTION OF AMYLOID- β PEPTIDE (x- \geq 41)

(57) Abstract

This invention provides methods of screening compounds for their ability to alter the production of $A\beta$ (x \geq 41) alone or in combination with $A\beta$ (x \leq 40). The methods involve administering compounds to cells, specifically measuring the amounts of $A\beta$ (x \leq 40) and $A\beta$ (x \geq 41) produced by the cells, and comparing these amounts to that produced by the cells without administration of the compounds.

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SCREENING COMPOUNDS FOR THE ABILITY TO ALTER THE PRODUCTION OF AMYLOID- β PEPTIDE ($x-\ge41$)

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BACKGROUND OF THE INVENTION

1. Field of the Invention

currently known.

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The present invention relates generally to neurology and, more particularly, to assays, such as immunoassays, for screening for compounds that specifically alter the production of various isoforms of $A\beta$.

Alzheimer's disease (AD) is a degenerative brain

disorder characterized clinically by progressive loss of memory, cognition, reasoning, judgment and emotional stability that gradually leads to profound mental deterioration and 25 ultimately death. AD is a very common cause of progressive mental failure (dementia) in aged humans and is believed to represent the fourth most common medical cause of death in the United States. AD has been observed in all races and ethnic groups worldwide and presents a major present and future public health problem. The disease is currently estimated to affect about two to three million individuals in the United States alone. AD is at present incurable. No treatment that effectively prevents AD or reverses its symptoms or course is

The brains of individuals with AD exhibit characteristic lesions termed senile plaques and neurofibrillary tangles. Large numbers of these lesions are generally found in patients with AD in several areas of the human brain important for memory and cognitive function.

Smaller numbers of these lesions in a more restricted anatomical distribution are sometimes found in the brains of aged humans who do not have clinical AD. Senile plaques and vascular amyloid deposits (amyloid angiopathy) also 5 characterize the brains of individuals beyond a certain age with Trisomy 21 (Down's Syndrome) and Hereditary Cerebral Hemorrhage with Amyloidosis of the Dutch-Type (HCHWA-D). principal chemical constituent of the senile plaques and vascular amyloid deposits characteristic of AD and the other 10 disorders mentioned above is a protein designated the amyloid-etapeptide (A β) or sometimes β AP, A β P or β /A4. A β was first purified and a partial amino acid sequence reported in Glenner and Wong (1984) Biochem. Biophys. Res. Commun. 120:885-890. The isolation procedure and the sequence data for the first 28 15 amino acids are described in U.S. Patent No. 4,666,829. of Aeta having amino acids beyond number 40 were first reported by Kang et al. (1987) Nature 325:733-736.

Roher et al. (1993) Proc. Natl. Acad. Sci. USA 90:10836-840 showed that $A\beta(1-42)$ is the major constituent in 20 neuritic plaques, including significant amounts of isomerized and racemized aspartyl residues as their NH_2 -termini. authors also reported that $A\beta(17-42)$ (p3(42)) predominates in diffuse (early) plaques, whereas $A\beta$ (1-40) is the major constituent in the meningeal vessel deposits, comprising 60% of 25 the total $A\beta$ in those vessels. Iwatsubo et al. (1994) Neuron 13:45-53 showed that $A\beta$ 42(43)-containing senile plaques are the major species of senile plaques in sporadic AD brains. Iwatsubo et al. (1995) Annals of Neurology 37:294-299 and Lemere et al. (1996) Neurobiology of Disease 3:16-32 reported 30 that $A\beta 42(43)$ is the major constituent of senile plaques in Down's syndrome brains and is the initially deposited Aetaspecies in the development of AD-type neuropathological legions in these patients. In addition, Gravina et al., (1995) J. Biol. Chem. 270:7013-7016 reported both biochemical and 35 immunocytochemical evidence that $A\beta42(43)$ peptides were the most abundant constituents of senile plaques in AD brains and exceeded the amounts of Aeta40 peptides in such plaques.

Molecular biological and protein chemical analyses conducted during the last several years have shown that ${\tt A}{\beta}$ is a small fragment of a much larger precursor protein, referred to as the β -amyloid precursor protein (APP), that is normally 5 produced by cells in many tissues of various animals, including humans. Knowledge of the structure of the gene encoding APP has demonstrated that Aeta arises as a peptide fragment that is cleaved from the carboxy-terminal end of APP by as-yet-unknown enzymes (proteases). The precise biochemical mechanism by 10 which the $A\beta$ fragment is cleaved from APP and subsequently deposited as amyloid plaques in the cerebral tissue and in the walls of cerebral and meningeal blood vessels is currently Importantly, Haass et al. (Nature 359:322-325) and Seubert et al. ((1992) Nature 359:325-327) discovered that 15 essentially all cells expressing the APP gene normally secrete an array of $A\beta$ peptides, and these peptides can readily be detected and assayed in cell culture fluid (conditioned media) and human biological fluids such as plasma and cerebrospinal fluid. It has subsequently been shown that these fluids 20 contain both the more abundant $A\beta 40$ -ending peptides and the less abundant $A\beta 42(43)$ -ending peptides (Dovey et al. (1993) Neuroreport 4:1039-1042 and Vigo-Pelfrey et al. (1993) J. Neurochem. 61:1965-68)

Several lines of evidence indicate that progressive

25 cerebral deposition of Aβ plays a seminal role in the
 pathogenesis of AD and can precede cognitive symptoms by years
 or decades (for review, see Schenk (1995) J. Med. Chem.
 38:4141-4154, Selkoe (1994) J. Neuropath. and Exp. Neurol.
 53:438-447 and Selkoe (1991) Neuron 6:487). One of the most

30 important lines of evidence is the discovery in 1991 that
 missense DNA mutations in the APP gene at amino acid 717 of the
 770-amino acid isoform of APP can be found in affected members
 but not unaffected members of several families with a
 genetically determined (familial) form of AD (Goate et al.

35 (1991) Nature 349:704-706; Chartier Harlan et al. (1991) Nature
 353:844-846; and Murrell et al. (1991) Science 254:97-99).
 Suzuki et al. (1994) "An increased percentage of long amyloid
 β-protein secreted by familial amyloid β-protein precursor

(βAPP717) mutants," Science 264:1336-1340 subsequently showed that, compared to normal individuals, the 717 mutation causes a higher relative production of the Aβ(1-42) peptide. In addition, a double mutation changing lysine⁶⁷⁰-methionine⁶⁷¹ to asparagine⁶⁷⁰-leucine⁶⁷¹ (with reference to the 770 isoform of APP) was reported in a Swedish family with familial AD in 1992 (Mullan et al. (1992) Nature Genet 1:345-347) and is referred to as the Swedish APP variant.

Genetic linkage analyses have demonstrated that the aforementioned mutations are the specific molecular cause of AD in the members of such families that carry these mutant APP genes. In addition, a mutation at amino acid 693 of the 770-amino acid isoform of APP has been identified as the cause of the A β deposition disease, Hereditary Cerebral Hemorrhage With Amyloidosis Dutch type (HCHWA-D), and a mutation from alanine to glycine at amino acid 692 appears to cause the phenotype of AD in some family members and the phenotype of HCHWA-D in others. The discovery of these APP mutations in genetically based cases of AD argues that genetic alteration of APP and subsequent deposition of its A β fragment can cause AD.

Recently, evidence has accumulated suggesting that $A\beta$ (42) plays the key role in the process of senile plague formation in AD. First, in vitro data demonstrate that $A\beta(42)$ accelerates the formation of $A\beta$ fibrils (and thus senile 25 plaques) by a nucleation dependent mechanism (Jarrett et al. (1993) Biochemistry 32:4693-4697). Second, while accounting for $\leq 10\%$ of total A β secreted from cells (roughly 90% is A β (40) (Dovey et al. (1993) Neuroreport 4:1039-1042; Asami-Odaka et al. (1995) "Long amyloid β -protein secreted from wild-type 30 human neuroblastoma IMR-32 cells." Biochemistry 34:10272-10278), $A\beta$ (42) is the major plaque component (Kang et al. (1987) Nature 325:733-736; Iwatsubo et al. (1994) Neuron 13:45-53; Iwatsubo et al. (1995) Ann. Neurol. 37:294-299; Gravina et al. (1995) J. Biol. Chem. 270:7013-7016; Lemere et al. (1996) 35 Neurobiology of Disease 3:16-32). Furthermore, all 3 early onset familial AD genes identified to date have been shown to lead to an increase in cellular secretion of $A\beta$ (42). Only the

Swedish APP missense mutation increases the secretion of both

 $A\beta$ (40) and $A\beta$ (42) peptides (Dovey et al. (1993) Neuroreport 4:1039-1042, whereas the APP717 mutations and the presentlin mutations appear not to increase $A\beta$ (40) peptides (Suzuki et al. (1994) Science 264:1336-1340; Scheuner et al. (1995) Neurosci. 5 Abstracts in press). Thus, the longer $A\beta(42)$ peptide appears to be a prime target for therapeutic intervention. none of the proteases involved in the major steps of APP processing have been definitively identified, including \gammasecretase, the protease which generates the C-terminus of $A\beta$. 10 It has generally been assumed that the same protease(s) generate both $A\beta(40)$ and $A\beta(42)$ and it has been shown that both forms share a common secretory mechanism which involves acidic intracellular compartments such as the late Golgi or early endosomes (Koo and Squazzo (1994) J. Biol. Chem. 269:17386-15 17389; Asami-Odaka et al. (1995) Biochemistry 34:10272-10278). Recently, Higaki et al. ((1995) Neuron, 14:651-659) have shown that the Calpain inhibitor, MDL 28170, inhibits the production of both total $A\beta$ and total p3 and leads to an accumulation of their respective 12 kDa and 10 kDa APP precursor fragments in 20 treated cells. These data suggest that the compound directly inhibits at least some form of γ -secretase although no data are provided as to what specific form of $A\beta$ and p3 are affected.

Despite the progress which has been made in understanding the underlying mechanisms of AD, there remains a need for assays to identify candidate compounds for preventing or treating the disease.

SUMMARY OF THE INVENTION

According to current theory, the processing of APP is believed to involve several specific cleavages by proteases. The enzyme that cleaves APP between amino acids 671/672 (referring to the β APP₇₇₀ isoform) is called β -secretase. The enzyme that cleaves between amino acids 687/688 of APP (16/17 of A β) is called α -secretase. Until now it was believed that cleavage of APP that yielded A β (40) and A β (42) was carried out by a single enzyme called γ -secretase. However, we have discovered that a compound can inhibit the production of A β (40) but not A β (42). In particular, we have discovered that

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compounds, thought to inhibit the production $A\beta$ in general, actually inhibit production of $A\beta(40)$ but not $A\beta(42)$. indicates that multiple γ -secretase mechanisms are at work which can be pharmacologically dissociated.

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Because $A\beta$ (42) is the major component of β -amyloid plaques and initiates amyloid plaque formation in AD patients, it is important to have tools to screen compounds to identify those that specifically inhibit the production of $A\beta$ (42) and $A\beta$ (40), either simultaneously or separately. The current 10 invention provides such assays.

This invention provides methods for determining whether a compound alters the amount of at least one $A\beta(x-\ge41)$ peptide produced by a cell. The methods involve administering the compound to a culture comprising the cell; measuring the 15 amount of the $A\beta(x-241)$ peptide, specifically, in a sample from the culture; and determining whether the measured amount is different than the amount expected in a sample from a culture comprising the cell to which no compound has been administered. A difference between the measured amount and the expected 20 amount indicates that the compound alters the amount of an $A\beta(x-\ge41)$ peptide produced by the cell.

In another aspect, this invention provides methods for determining whether a compound alters the amount of at least one $A\beta(x-\ge41)$ peptide produced by a cell and alters the 25 amount of either total $A\beta$ or at least one $A\beta(x-\leq 40)$ peptide produced by the cell. The methods involve administering the compound to a culture comprising the cell; measuring the amount of the $A\beta(x-241)$ peptide, specifically, in a sample from the culture; measuring the amount of total $A\beta$ or the $A\beta(x-\le 40)$ 30 peptide, specifically, in a sample from the culture; and determining whether the measured amounts are different than the amounts expected in a sample from a culture comprising the cell to which no compound has been administered. Differences between the measured amounts and the expected amounts indicate 35 that the compound alters the amount of the $A\beta(x-\ge41)$ peptide by a cell and/or the amount of total $A\beta$ or the $A\beta$ (x- \leq 40) peptide by the cell.

In one embodiment, the amount of the $A\beta$ peptides are measured by immunoassay and, in particular, sandwich immunoassay comprising capture binding substances bound to a solid phase and a labeled detection binding substance.

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In sandwich assays for determining the amount of at least one $A\beta(x-\ge41)$ peptide, the capture antibody preferably is specific for $A\beta(x-\ge 41)$ peptides, e.g., raised against peptide NH2-Cys-NH-CH2-(CH2)5-CO-GLMVGGVVIA-COOH (SEQ ID NO:4). detection binding substance in this assay can be an antibody 10 specific for $A\beta$ peptides whose amino-terminal amino acid is no. 1 of $A\beta$, or can be specific for an epitope within the junction region of $A\beta$. In another embodiment the capture binding substance for measuring the amount of at least one of $A\beta(x-241)$ peptide is specific for an epitope within the junction region 15 of $A\beta$ and the detection binding substance is an antibody specific for $A\beta(x-\ge 41)$.

In sandwich assays for determining the amount of at least one $A\beta(x-40)$ peptide, the capture binding substance preferably is an antibody specific for Aβ(x-≤40) peptides, 20 e.g., raised against the peptide NH₂-Cys-NH-CH₂-(CH₂)₅-CO-GLMVGGVV-COOH (SEQ ID NO:5). The labeled detection binding substance can be an antibody specific for the $A\beta$ peptides whose amino-terminal amino acid is no. 1 of $A\beta$ or an antibody specific for an epitope within the junction region of $A\beta$. 25 another embodiment the capture binding substance for measuring the amount of at least one of $A\beta(x-\le 40)$ peptide is specific for an epitope within the junction region of $A\beta$ and the detection binding substance is an antibody specific for $A\beta(x-\le 40)$.

In sandwich assays for determining the amount of 30 total $A\beta$, the capture binding substance preferably is an antibody specific for an epitope within the junction region of $A\beta$. The detection binding substance preferably is specific for $A\beta$ peptides whose amino-terminal amino acid is no. 1 of $A\beta$.

In another embodiment of an immunoassay, the step of 35 measuring the amount of the $A\beta(x-\ge41)$ peptide, total $A\beta$ or the $A\beta(x-\leq 40)$ peptide in a sample from the culture comprises: pulsing the culture with a radioactive label for protein; chasing the culture without a radioactive label; administering

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the compound to the cell during the chase period; contacting a sample from the culture with a binding substance specific for $A\beta(x-\ge41)$ peptides; contacting a sample from the culture with a binding substance specific for total $A\beta$ or $A\beta(x-\le40)$ peptide; and determining the amount of radioactive label attached to the binding substances.

In other embodiments of the methods the culture comprises primary human neurons, primary neurons from a transgenic PDAPP mouse (i.e., a transgenic mouse whose cells harbor a PDAPP construct), a 293 human kidney cell line, a human neuroglioma cell line, a human HeLa cell line, a primary endothelial cell line, a primary human fibroblast line, a primary lymphoblast line, human mixed brain cells, or a Chinese hamster ovary (CHO) cell line. In one embodiment the cell is a host cell transfected with a recombinant expression vector encoding a human APP, e.g., a Hardy mutation such as V717F or the Swedish mutant; causing the cell to overproduce Aβ(x-≥41) peptides. In another aspect, the methods further comprise the step of determining whether the compound is toxic to the cell.

In another aspect this invention provides kits for specifically detecting at least one $A\beta(x-\ge 41)$ peptide and at least one $A\beta(x-\le 40)$ peptide in a sample. The kits include a binding substance specific for $A\beta(x-\ge 41)$ peptides; and a binding substance specific for $A\beta(x-\le 40)$ peptides.

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In another aspect this invention provides kits for specifically detecting at least one $A\beta(x-\ge41)$ peptide and either total $A\beta$ or at least one $A\beta(x-\le40)$ peptide in a sample in a sandwich immunoassay. The kits include at least two different binding substances for measuring the amount of $A\beta(x-30)$ peptide; and at least two different binding substances for measuring the amount of total $A\beta$ or $A\beta(x-\le40)$ peptides.

In another aspect this invention provides methods for determining whether a compound alters the amount of at least one $A\beta(x-\ge41)$ peptide produced by a non-human mammal and alters the amount of either total $A\beta$ or at least one $A\beta(x-\le40)$ peptide produced in the non-human mammal. The methods involve measuring a first amount of the $A\beta(x-\ge41)$ peptide in a sample from a non-human animal used as a model of Alzheimer's disease;

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measuring a first amount of total $A\beta$ or the $A\beta(x-40)$ peptide in a sample from the non-human animal; administering the compound to the non-human animal; measuring a second amount of the $A\beta(x-41)$ peptide in a sample from the non-human animal; measuring a second amount of total $A\beta$ or the $A\beta(x-40)$ peptide in a sample from the non-human animal; and comparing the first amounts with the second amounts. The comparison indicates whether the compound increases, decreases, or leaves unchanged the amount of the $A\beta(x-41)$ peptide and increases, decreases, or leaves unchanged the amount of the $A\beta(x-41)$ peptide and increases, decreases, or leaves unchanged the amount of the $A\beta(x-40)$ peptide. In certain embodiments the non-human animal is a rodent, in particular, a mouse. The non-human animal can harbor a copy of an expressible transgene sequence which encodes a Hardy mutation (e.g., V717F) or the Swedish mutation of human β -amyloid precursor protein (APP).

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1. Shows standard curves of $A\beta$ (1-40) and $A\beta$ (1-42) with 2G3 as a capture antibody.

Fig. 2. Shows standard curves of $A\beta$ (1-40) and $A\beta$ (1-42) with 21F12 as a capture antibody.

Fig. 3. Shows that compound MDL 28170 influences β APP metabolism. (0) untreated K695sw cells; (200) K695sw cells treated with 200 μ M MDL 28170 during the chase period.

25 Lanes 1,2: aliquot of total chase media run directly on the gel. Lanes 3,4: antibody 1736 immunoprecipitation of chase media. Lanes 5,6: antibody 192sw immunoprecipitation of chase media. Note that β -cut APPs (lanes 5,6) runs slightly below α -cut APPs (lanes 3,4), as expected. Lanes 7,8: antibody C7 immunoprecipitation of cell lysates. Lanes 9,10: antibody 1282 immunoprecipitation of cell lysates.

Figs. 4A-4B. Differential inhibition of $A\beta$ (42) and $A\beta$ (40) formation. (A) Labeled K695sw cells were chased with the indicated concentrations of MDL 28170, and the chase media were precipitated with 21F12 (upper panel) followed by antibody 1282 (lower panel). (B) Quantitation of the effect of 200 μ M MDL 28170 on $A\beta$ and β 3 in the chase media by phosphorimaging. The bars show the relative pixel number compared to an

untreated control (set at 100%). The decreases in $A\beta_{total}$ and $p3_{total}$ relative to an untreated control were significant (*) (two-tailed t-test, n=4, p<0.001), whereas the decreases in $A\beta$ (42) and p3(42) upon treatment with MDL 28170 did not reach significance. The difference in reduction of $A\beta$ (42) level vs. total $A\beta$ level is significant (two-tailed t-test, n=4, p<0.01). The difference in inhibition of p3(42) vs. total p3 is also significant (two-tailed t-test, n=4, p<0.05).

Figs. 5A-5F. Differential inhibition of AB 10 production by K695sw cells under a variety of conditions. (A) treatment with 1 μ M PDBu (P) decreases both A β (42) and A β _{total}. Upper panel: antibody 21F12 precipitation, lower panel: subsequent antibody 1282 precipitation. (B-E) K695sw cells were chased with (M) or without (0) 200 μM MDL 28170, and the 15 chase media were immunoprecipitated (B) first with antibody BC05 (upper panel) and then with antibody 1282 (lower panel); (C) first with antibody C42 (upper panel) and then with antibody 1282 (lower panel); (D) first with antibody 2G3 (upper panel) and then with antibody 21F12 (lower panel); (E) after 20 splitting the medium into two aliquots, either with antibody 21F12 (upper panel) or antibody 1282 (lower panel). (F) cells were labeled for 3 h in the presence of 100 μM MDL 28170, and the medium was immunoprecipitated first with antibody 21F12 (upper panel) and then with antibody 1282 (lower panel).

Figs. 6A-6C. Differential inhibition of Aβ production in different cell types. 35S-methionine labeled cells were chased with the indicated concentrations of MDL 28170 and precipitated with antibody 21F12 (upper panel) followed by antibody 1282 (lower panel). (A) V717F cells. The relatively low APP expression leads to a faint Aβ42 band. (B) CHO695 cells. Note that in CHO cells the p3 brands migrate as doublets, as described (Koo and Squazzo, (1994) "Evidence that production and release of amyloid β-protein involves the endocytic pathway." J. Biol. Chem. 269:17386-17389). (C) SKN695 cells. While Aβ(42) and p3(42) are slightly increased at 200 μM MDL 28170, Aβtotal and p3total are decreased.

Fig. 7. Shows a summary sketch of some of the known proteolytic processing pathways of APP.

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Fig. 8A. Shows a schematic of a combination cDNA/genomic coding sequence allowing alternative splicing of the KPI and OX-2 exons.

Fig. 8B. Shows a schematic of a combination 5 cDNA/genomic coding sequence bearing a mutation at position 717 and allowing alternative splicing of the KPI and OX-2 exons.

Fig. 9. Diagram of the intermediate constructs used to construct the APP splicing cassette and the PDAPP vector.

Fig. 10A-D. The DNA sequence (SEQ ID NO:2) and the 10 deduced amino acid sequence (SEQ ID NO:3) of APP770.

Fig. 11. Diagram of the genomic region of APP present in the PDAPP construct. The sizes of original introns 6, 7 and 8, as well as the sizes of the final introns are indicated in the diagram. The locations of the deletions in introns 6 and 8 present in the PDAPP construct also are indicated.

Fig. 12. Schematic map of the PDAPP vector, a combination cDNA/genomic APP construct.

Figs. 13A-13O depict the percent inhibition of $A\beta$ by several compounds using the brain cell culture method described in Example VIII. The charts show inhibition of total $A\beta$ ("% $A\beta$ inhibition"), inhibition of $A\beta$ (42) ("% 1-42 inhibition") and inhibition of the metabolism of MTT ("% MTT inhibition") (greater inhibition indicates greater cytotoxicity).

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DETAILED DESCRIPTION OF THE INVENTION

I. <u>DEFINITIONS</u>

The term "binding substance" refers to a polypeptide substantially encoded by an immunoglobulin gene or immunoglobulin genes, or fragments thereof, which specifically bind and recognize an analyte (antigen). The recognized immunoglobulin genes include the kappa, lambda, alpha, gamma, delta, epsilon and mu constant region genes, as well as the myriad immunoglobulin variable region genes. Antibodies exist, e.g., as intact immunoglobulins or as a number of well characterized fragments produced by digestion with various peptidases. This includes, e.g., Fab' and F(ab)', fragments. The term "binding substance," as used herein, also includes

antibody fragments either produced by the modification of whole antibodies or those synthesized *de novo* using recombinant DNA methodologies.

The term "immunoassay" is an assay that utilizes a binding substance to specifically bind an analyte. The immunoassay is characterized by the use of specific binding properties of a particular antibody to isolate, target, and/or quantify the analyte.

A binding substance "specifically binds to" or "is 10 specifically immunoreactive with" a protein when the binding substance functions in a binding reaction which is determinative of the presence of the protein in the presence of a heterogeneous population of proteins and other biologics. Thus, under designated immunoassay conditions, the specified 15 binding substances bind preferentially to a particular protein and do not bind in a significant amount to other proteins present in the sample. Specific binding to a protein under such conditions requires an antibody that is selected for its specificity for a particular protein. A variety of immunoassay 20 formats may be used to select binding substances specifically immunoreactive with a particular protein. For example, solid-phase ELISA immunoassays are routinely used to select monoclonal antibodies specifically immunoreactive with a protein. See Harlow and Lane (1988) Antibodies, A Laboratory 25 Manual, Cold Spring Harbor Publications, New York, for a description of immunoassay formats and conditions that can be used to determine specific immunoreactivity.

A "label" is a composition detectable by spectroscopic, photochemical, biochemical, immunochemical, or chemical means. For example, useful labels include 32p, fluorescent dyes, electron-dense reagents, enzymes (e.g., as commonly used in an ELISA), biotin, dioxigenin, or haptens and proteins for which antisera or monoclonal antibodies are available. Binding substances can be made detectible, e.g., by incorporating a radio-label into the peptide, and used to detect antibodies specifically reactive with the peptide. A label often generates a measurable signal, such as

radioactivity, fluorescent light or enzyme activity, which can be used to quantitate the amount of bound label.

II. <u>IN VITRO SCREENING</u>

5 Alzheimer's disease is characterized by the initial deposition of $A\beta(x-241)$ in the form of amyloid plagues in the Therefore, effective treatments for AD are expected to decrease the production of these peptides, whereas agents that hasten progress of the disease are expected to increase 10 production of the peptide. Prior screening methods looked for compounds that decreased total A β . However, since A β (x- \geq 41) peptides are a small fraction of total $A\beta$, those assays could not determine whether the compound specifically inhibited $A\beta$ (x-≥41) peptides. As results described herein indicate, compounds 15 can alter the production of $A\beta(x-\leq 40)$ but not $A\beta(x-\geq 41)$. Because $A\beta$ (42) is the major constituent of neuritic plaques, it is useful to identify compounds that specifically inhibit the production of $A\beta(x-\ge41)$ peptides, either in addition to, or instead of $A\beta(x-\leq 40)$ peptides. Accordingly, this invention 20 provides methods for screening compounds that specifically elevate or decrease the production of the amount of $A\beta(x-241)$ by a cell and compounds that elevate or decrease production of both $A\beta(x-241)$ and $A\beta(x-440)$ (e.g., total $A\beta$), or of one or the other of these peptides. Compounds that decrease production of 25 $A\beta(x-241)$ are candidates for use in treating the disease, while compounds that increase its production may hasten the disease and are to be avoided by humans.

Screening methods of this invention for determining whether a test compound specifically alters the amount of $A\beta$ (x-30 \ge 41) produced by a cell involve administering the compound to the cell, usually in culture, measuring the amount of $A\beta$ (x- \ge 41) specifically produced by the cell, and determining whether this amount is greater than, less than or the same as an amount the cell is expected to produce in the absence of the compound. If the amounts are different, then the compound affects the production of $A\beta$ (x- \ge 41) by the cell. This amount can be measured, for example, in a sample from the culture, such as

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medium conditioned by the cell in culture, or in extracts derived from cells harvested from the culture.

The expected amount generally will be a control amount determined by measuring $A\beta(x-\ge 41)$ produced by the cell in the absence of the compound. However, one also may determine the expected amount by extrapolation; measuring the amount of $A\beta(x-\ge 41)$ produced upon administration of different amounts of the compound to the cell, and using these figures to calculate the expected amount. In certain instances measuring a control amount for the purposes of comparison may not be necessary because the effect of the compound on $A\beta(x-\ge 41)$ production is clear. For example, a compound may render $A\beta(x-\ge 41)$ undetectable in a cell that normally produces detectable amounts, indicating that the compound decreases $A\beta(x-\ge 41)$ production from the amount expected in its absence.

In another aspect, this invention provides screening methods for determining whether a compound alters the production of $A\beta(x-\ge 41)$ by a cell to a different degree than it alters the production of total $A\beta$ or $A\beta(x-\le 40)$ by the cell.

20 These methods are useful for determining whether a compound alters the production of $A\beta(x-\ge 41)$ in addition to total $A\beta$, or alters the production of one or the other of $A\beta(x-\ge 41)$ and $A\beta(x-\le 40)$. The methods involve administering the compound to the cell (usually in culture). Then, the degree to which the compound alters the production of $A\beta(x-\ge 41)$, specifically, by the cell is determined. The degree to which the compound alters the production of total $A\beta$ or $A\beta(x-\le 40)$ by the cell also is determined. Then, the degrees are compared. The comparison indicates whether the compound alters the production of $A\beta(x-\le 41)$ instead of or in addition to $A\beta(x-\le 40)$.

Determining the degree to which a compound alters the production of one or the other peptide generally involves measuring the specific amount of the peptide in a sample from the culture; and comparing it with the amounts expected in a sample from a culture comprising the cell to which no compound has been administered.

III. AMYLOID- β PEPTIDE AND RELATED PROTEINS AND PEPTIDES

Various cellular processing pathways for APP are presented in Fig. 7. The terms "amyloid- β peptide," "A β " or " β AP" as used herein refer to an approximately 4.2 kD protein 5 which, in the brains of AD, Down's Syndrome, HCHWA-D and some normal aged subjects, forms the subunit of the amyloid filaments comprising the senile (amyloid) plaques and the amyloid deposits in small cerebral and meningeal blood vessels (amyloid angiopathy). A β can occur in a filamentous polymeric 10 form (in this form, it exhibits the Congo-red and thioflavin-S dye-binding characteristics of amyloid described in connection therewith). A β can also occur in a non-filamentous form ("preamyloid" or "amorphous" or "diffuse" deposits) in brain tissue, in which form no birefringent staining by Congo red 15 occurs. A portion of this protein in the insoluble form obtained from meningeal blood vessels is described in U.S. Patent No. 4,666,829.

 $A\beta$ is an approximately 39-43 amino acid fragment of a large membrane-spanning glycoprotein, referred to as the β 20 amyloid precursor protein (APP), encoded by a gene on the long arm of human chromosome 21. Forms of $A\beta$ longer than 43 amino acids are also contemplated herein. $A\beta$ is further characterized by its relative mobility in SDS-polyacrylamide gel electrophoresis or in high performance liquid
25 chromatography (HPLC). A sequence for a 43-amino acid-version of $A\beta$ is:

1 Asp Ala Glu Phe Arg His Asp Ser Gly Tyr

30 <u>11</u> Glu Val His His Gln Lys Leu Val Phe Phe

21
Ala Glu Asp Val Gly Ser Asn Lys Gly Ala

31
Ile Ile Gly Leu Met Val Gly Gly Val Val

40 Ile Ala Thr (SEQ ID NO:1).

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As used herein, $A\beta$ also refers to related polymorphic forms of $A\beta$, including those that result from mutations in the $A\beta$ region of the APP gene.

The term "A β fragment" as used herein refers to fragments and degradation products of A β which are generated at low concentrations by mammalian cells. Particular A β fragments have a molecular weight of approximately 3 kD and are presently believed to include peptides with, for example, amino acid residues 3-34, 6-27, 6-34, 6-35, 6-42, 11-34, 11-40, 11-43, 12-43, 17-40 and 17-42 of A β (Vigo-Pelfrey et al. (1993) J. Neurochem. 61:1965-1968).

As used herein, the term "A β (x- \geq 41)" refers to A β or an A β fragment whose amino-terminus begins at amino acid number 1 of A β or thereafter (i.e., which is amino-terminally truncated), and whose carboxy-terminus extends beyond amino acid number 40. These peptides and fragments comprise a heterogenous group. For example, A β (6-42), A β (11-43) and 15 A β (12-43) all have been found in the CSF. However, this list is not meant to be exclusive. Other peptides from among the group are presumed to exist in the culture media of cells expressing APP and are detectable with the methods described herein. As used herein the term "A β (42)" refers to A β or an A β fragment whose C-terminal amino acid is # 42 of A β .

As used herein, the term "A β (x- \leq 40)" refers to A β or an A β fragment whose amino-terminus begins at amino acid number 1 of A β or which is amino-terminally truncated, and whose carboxy-terminus extends no further than amino acid number 40. These peptides and fragments also comprise a heterogenous group. The term "A β (40)" refers to A β or an A β fragment whose C-terminal amino acid is # 40 of A β .

As used herein, the term "p3" refers to a peptide whose amino acid sequence is substantially similar to $A\beta$, but whose amino-terminal amino acid begins at amino acid 17 of $A\beta$. The term "p3 fragment" as used herein refers to fragments and degradation products of p3. Whereas p3 is produced through a different processing pathway than $A\beta$, for the purposes of the detection methods of this invention, p3 and p3 fragments are considered to be a subset of $A\beta$ peptides, because certain detection techniques that recognize $A\beta$ solely from the carboxy-terminus generally also will recognize p3. Also it is shown

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that the same apparent mechanisms generate the p3 and ${\rm A}\beta$ carboxy-termini.

The term "A β junction region" as used herein refers to a region of A β which is centered at the site between amino acid residues 16 and 17 (Lys¹6 and Leu¹7), which is a principal target for proteolytic processing of APP. Such processing, referred to as " α -secretory" processing, results in a variety of APP fragments which may, for example, terminate at amino acid 16 of A β and which, therefore, are potentially

immunologically cross-reactive with antibodies to the intact $A\beta$ molecule which are to be used in the methods of the present invention. Antibodies raised against a synthetic peptide including amino acid residues 13-28 have been found to display the requisite specificity for the junction region.

The term "amyloid- β precursor protein" (APP) as used herein is defined as a polypeptide that is encoded by a gene of the same name localized in humans on the long arm of chromosome 21 and that includes the A β region within the carboxyl third of its coding region. APP is a glycosylated, single-membrane-

spanning protein expressed in a wide variety of cells in many mammalian tissues. Examples of specific isotypes of APP which are currently known to exist in humans are the 695-amino acid polypeptide described by Kang et al. (1987) Nature 325:733-736; the 751-amino acid polypeptide described by Ponte et al. (1988)

Nature 331:525-527 (1988) and Tanzi et al. (1988) Nature 331:528-530; and the 770-amino acid polypeptide described by Kitaguchi et al. (1988) Nature 331:530-532. Examples of specific variants of APP include point mutations which can differ in both position and resultant neuropathological

30 phenotype (for review of known variant mutations see Hardy (1992) Nature Genet. 1:233-234).

The term "A β -related condition" as used herein is defined as including Alzheimer's disease (which includes familial Alzheimer's disease), Down's Syndrome, HCHWA-D, and advanced aging of the brain.

IV. CELLS EXPRESSING $A\beta(x-\ge 41)$

Because it is a therapeutic goal to discover compounds that inhibit the production of $A\beta(x-\ge41)$, the test cells used in the methods of this invention generally are ones that are able to secrete $A\beta(x-\ge41)$.

In vitro monitoring of $A\beta(x-\ge41)$ levels in conditioned medium from a suitable cell culture may be used for drug screening. By growing cells under conditions which result in the secretion of $A\beta(x-\ge41)$ into the culture medium, and exposing the cells to test compounds, the effect of these test compounds on $A\beta(x-\ge41)$ secretion can be observed.

Suitable cell lines include human and animal cell lines, such as, preferably, primary human neurons, and primary neurons from transgenic mice harboring human APP genes, e.g., 15 cells from a transgenic PDAPP animal (e.g., mouse), as well as a 293 human kidney cell line, a human neuroglioma cell line, a human HeLa cell line, a primary endothelial cell line (e.g., HUVEC cells), a primary human fibroblast line or a primary lymphoblast line (including endogenous cells derived from 20 patients with APP mutations), a primary human mixed brain cell culture (including neurons, astrocytes and neuroglia), or a Chinese hamster ovary (CHO) cell line. Particularly useful are cells stably transfected with APP₆₉₅ carrying the mutation V717I (valine to isoleucine at position 717 in the APP770 numbering 25 system). Cell lines which preferentially increase the levels or ratios of $A\beta(x-\ge41)$ would be particularly useful in the methods of invention. Useful mutants at position 717 (the Hardy mutation) include V717F, V717I or V717G.

Preferred for use in drug screening methods according to the present invention are cell lines capable of expressing APP variants which overproduce A\$\beta\$. By "overproduce," it is meant that the amount of A\$\beta\$ produced from the variant APP will be at least about one-and-a-half times and preferably at least two or five times greater than the amount produced from any or all of the normal APP isoforms, e.g., the 695, 751, and 770 amino acid isoforms which have been previously described. Particularly preferred are APP variants having one or several amino acid substitutions directly amino-terminal of the A\$\beta\$

cleavage site. For example, K293 cells which express an APP_{695} DNA bearing a double mutation (Lys^{595} -> Asn^{595} and Met^{596} -> Leu^{596} (695 numbering system)) found in a Swedish FAD family produce approximately five-to-eight-fold more $A\beta$ than cells expressing normal APP (Citron et al. (1992) Nature, 360:672-674). The mutation at residue 596 appears to be principally responsible for the increase.

V. <u>EXPRESSION VECTORS FOR APP</u>

Host cells transfected with a recombinant expression vector that encodes APP also are useful as cells in the screening methods of this invention. A plasmid that carries sequences encoding APP is pCMV695 (Selkoe et al. (1988) Proc. Natl. Acad. Sci USA 85:7341-7345).

Nucleic acids encoding APP can be obtained by methods known in the art. For example, a nucleic acid encoding an APP can be isolated by polymerase chain reaction of cDNA or genomic DNA from a human brain cDNA library or a human genomic library using primers based on the DNA sequence of APP. PCR methods are described in, for example, U.S. Pat. No. 4,683,195; Mullis et al. (1987) Cold Spring Harbor Symp. Quant. Biol. 51:263; and Erlich, ed., PCR Technology, (Stockton Press, NY, 1989).

Mutant versions of APP, such as the Swedish mutation, can be made by site-specific mutagenesis of other nucleic acids encoding APP, or by random mutagenesis caused by increasing the error rate of PCR of the original polynucleotide with 0.1 mM MnCl₂ and unbalanced nucleotide concentrations.

The construction of expression vectors and the expression of genes in transfected cells involves the use of molecular cloning techniques also well known in the art.

Sambrook et al., Molecular Cloning -- A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, (1989) and Current Protocols in Molecular Biology, F.M. Ausubel et al., eds., (Current Protocols, a joint venture between Greene Publishing Associates, Inc. and John Wiley & Sons, Inc.).

Nucleic acids used to transfect cells with sequences coding for expression of the polypeptide of interest generally will be in the form of an expression vector that includes

expression control sequences operatively linked to a nucleotide sequence coding for expression of the polypeptide. As used herein, the term nucleotide sequence "coding for expression of" a polypeptide refers to a sequence that, upon transcription and 5 translation of mRNA, produces the polypeptide. As any person skilled in the art recognizes, this includes all degenerate nucleic acid sequences encoding the same amino acid sequence. This can include sequences containing, e.g., introns. As used herein, the term "expression control sequences" refers to 10 nucleic acid sequences that regulate the expression of a nucleic acid sequence to which it is operatively linked. Expression control sequences are "operatively linked" to a nucleic acid sequence when the expression control sequences control and regulate the transcription and, as appropriate, 15 translation of the nucleic acid sequence. Thus, expression control sequences can include appropriate promoters, enhancers, transcription terminators, a start codon (i.e., ATG) in front of a protein-encoding gene, splicing signals for introns, maintenance of the correct reading frame of that gene to permit 20 proper translation of the mRNA, and stop codons.

The recombinant nucleic acid can be incorporated into an expression vector comprising expression control sequences operatively linked to the recombinant nucleic acid. The expression vector can be adapted for function in prokaryotes or eukaryotes by inclusion of appropriate promoters, replication sequences, markers, etc.

The expression vector can be transfected into a host cell for expression of the recombinant nucleic acid. Host cells can be selected for high levels of expression in order to purify the protein. The host cell can be a prokaryotic or eukaryotic cell selected to study the activity of an enzyme produced by the cell. The cell can be, e.g., a cultured cell or a cell in vivo.

Transfected host cells useful in this invention

35 include human kidney 293 cell lines such as K695sw, and K695⁷¹⁷¹,

glioma cell lines such as HS695, and neuroblastoma cell lines
such as SKN695, described in the Experimental section.

VI. MEASURING $A\beta(x-\ge 41)$

 $A\beta$ peptides can be detected by any method known in the art. Preferably, the method involves an immunoassay employing binding substances specific for the peptides. Optionally, one can detect $A\beta$ peptides by determining their size, e.g., by HPLC or by mass spectrometry.

A. Binding Substances

One step of the screening methods of this invention involves measuring the amount of at least one $A\beta(x-\ge 41)$ 10 peptide, specifically, in a sample. Measuring $A\beta(x-\ge 41)$ peptides specifically means measuring $A\beta(x-\ge 41)$ peptides so as to distinguish that molecule from shorter species of $A\beta$, i.e., those species whose carboxy-terminus extends no further than amino acid # 40 of $A\beta$.

Specific measurement of $A\beta(x-\ge41)$ preferably is performed by the use of binding substances that specifically recognize $A\beta(x-\ge41)$ peptides, e.g., binding substances that recognize amino acids of $A\beta$ beyond amino acid # 40.

Another method of this invention involves screening compounds to determine their ability to alter the production of both $A\beta(x-\ge41)$ peptides and total $A\beta$ or $A\beta(x-\le40)$ peptides. Such methods can involve the use of binding substances that can distinguish $A\beta(x-\le40)$ peptides from longer species of $A\beta$, such as $A\beta(x-\ge41)$ peptides.

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B. <u>Immunoassays</u>

The use of immunological detection techniques, i.e., immunoassays employing binding substances, is preferred. Particularly suitable detection techniques include ELISA,

30 Western blotting, radioimmunoassay, and the like. Suitable immunological methods employing a single antibody are also contemplated, for example, radioimmunoassay using an antibody specific for ≥41 forms of Aβ, or single antibody ELISA methods. It will be clear that the particular forms of Aβ detected by such methods depend upon the particular binding substances employed. For example, binding substances directed to the junction region may detect Aβ(x-≥41) peptides whose amino termini do not extend to amino acid no. 1 of Aβ. Also, binding

substances directed to the carboxy-terminal end of $A\beta(x-\ge 41)$ may detect peptides ending at amino acids 41, 42 or 43. Therefore, determining the specificity of the binding substances will assist in determining exactly which $A\beta(x-\ge 41)$ peptides the assay is detecting.

In one embodiment, the method to detect $A\beta(x-\ge41)$ peptides is an immunoassay involving two antibodies. One antibody is specific for an epitope containing amino acids beyond number 40 in $A\beta$, and another antibody is capable of distinguishing $A\beta$ and $A\beta$ fragments from other APP fragments which might be found in the sample. In particular, it has been found that antibodies which are monospecific for the junction region of $A\beta$ are capable of distinguishing $A\beta$ from other APP fragments. The junction region of $A\beta$ is centered at amino acid residues 16 and 17, typically spanning amino acid residues ~13 to ~28. Such "junction-recognizing" antibodies may be prepared using synthetic peptides having that sequence as an immunogen.

"sandwich" assay. This assay involves a capture binding substance, usually bound to a solid phase, and a labelled detection binding substance. In this method, $A\beta(x-\ge 41)$ peptides are captured from the sample using a first binding substance specific for $A\beta(x-\ge 41)$ peptides (usually bound to a solid phase). The capture of $A\beta(x-\ge 41)$ peptides is detected using a labeled second binding substance specific for $A\beta$. Labeled binding substances include, for example, those directed to the junction region (amino acids ~13 to ~28) or binding substances specific for amino-terminal amino acids (1-5 or 1-12).

Particular methods for preparing such antibodies and utilizing such antibodies in an exemplary ELISA are set forth in the Experimental section hereinafter and in related United States patent application 07/965,972, supra. A sandwich assay using an antibody against the junction region can be used to specifically measure $A\beta$ and $A\beta$ fragments whose amino-terminus begins before amino acid 13 of $A\beta$. Such assays do not recognize p3 or p3 fragments, since those peptides begin at amino acid # 17 of $A\beta$.

Antibodies specific for $A\beta(x-z41)$, i.e., which do not cross react with $A\beta(z40)$, are particularly useful in the methods of this invention. These antibodies can be made by immunizing animals with synthetic peptides that include amino acids beyond number 40 of $A\beta$. For example, the synthetic peptide can include amino acids 33-42. A specific example of the production of such an antibody is provided in the Experimental section.

The particular peptides measured from among the group 10 of all $A\beta(x-241)$ depends on the particular measuring method In the case of using binding substances, such as antibodies, the binding substance can be directed to one or more from among the group of peptides. For example, an antibody raised against amino acids 33-42 of Aeta that does not 15 cross react with $A\beta(1-40)$ will bind to $A\beta(x-42)$. It also may bind to $A\beta(x-41)$ and $A\beta(x-43)$. According to one embodiment of the invention, the method involves determining the amount of $A\beta(x-\ge41)$ having at least amino acids 13-41 of $A\beta$. species can be measured using a sandwich assay employing 20 antibodies that recognize the junction region (amino acids 13-26) and antibodies produced by immunization with a hapten having $A\beta$ amino acids 33-42. Total $A\beta$ can be measured using a capture antibody to the junction region (e.g., the 266 antibody, described herein) and a reporter antibody that should 25 detect virtually all the $A\beta$ peptides and $A\beta$ fragments, e.g., an antibody raised against amino acids 1-12 of $A\beta$.

C. <u>Pulse-Chase Assays</u>

Another method of measuring the amount of $A\beta(x-\ge41)$ in a sample involves pulse-chase procedures. In these methods, the culture is pulsed with a radioactive label for protein, e.g., a radioactive amino acid such as ^{35}S methionine. Then the culture is chased without the label. Then the compound is administered to the cells. Then the cells are contacted with a binding substance specific for $A\beta(x-\ge41)$ and the amount of radioactive label attached to the binding substance is determined.

D. Preparation of Antibodies

Antibodies specific for $A\beta$ can be prepared, e.g., by immunizing an animal with a peptide whose amino acid sequence corresponds with amino acids ~13 to ~28 of $A\beta$. Antibodies specific for $A\beta(x-241)$ can be prepared, e.g., by immunizing an animal with a peptide whose amino acid sequence corresponds with amino acids 33-42 of $A\beta$. Antibodies specific for $A\beta(x-40)$ can be prepared, e.g., by immunizing an animal with a peptide whose amino acid sequence corresponds with amino acids 33-40 or 28-40 of $A\beta$. Antibodies against the junction region are useful for detecting total $A\beta$.

Synthetic polypeptide haptens may be produced by the well-known Merrifield solid-phase synthesis technique in which amino acids are sequentially added to a growing chain (Merrifield (1963) J. Am. Chem. Soc. 85:2149-2156). Suitable peptide haptens will usually comprise at least five contiguous residues within $A\beta$ and may include more than six residues. The amino acid sequences may be based on the sequence of $A\beta$ set forth above.

Once a sufficient quantity of polypeptide hapten has been obtained, it may be conjugated to a suitable immunogenic carrier, such as serum albumin, keyhole limpet hemocyanin, or other suitable protein carriers, as generally described in Hudson and Hay, Practical Immunology, Blackwell Scientific Publications, Oxford, Chapter 1.3, 1980. An exemplary immunogenic carrier utilized in the examples provided below is α-CD3ε antibody (Boehringer-Mannheim, Clone No. 145-2C11).

Antibodies specific for the desired epitope may be produced by in vitro or in vivo techniques. In vitro

30 techniques involve exposure of lymphocytes to the immunogens, while in vivo techniques require the injection of the immunogens into a suitable vertebrate host. Suitable vertebrate hosts are non-human, including mice, rats, rabbits, sheep, goats, and the like. Immunogens are injected into the animal according to a predetermined schedule, and the animals are periodically bled, with successive bleeds having improved titer and specificity. The injections may be made intramuscularly, intraperitoneally, subcutaneously, or the

like, and an adjuvant, such as incomplete Freund's adjuvant, may be employed.

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If desired, monoclonal antibodies can be obtained by preparing immortalized cell lines capable of producing 5 antibodies having desired specificity. Such immortalized cell lines may be produced in a variety of ways. Conveniently, a small vertebrate, such as a mouse, is hyperimmunized with the desired immunogen by the method just described. The vertebrate is then killed, usually several days after the final 10 immunization, the spleen cells removed, and the spleen cells The manner of immortalization is not critical. immortalized. Presently, the most common technique is fusion with a myeloma cell fusion partner, as first described by Kohler and Milstein (1975) Nature 256:495-497. Other techniques including EBV 15 transformation, transformation with bare DNA, e.g., oncogenes, retroviruses, etc., or any other method which provides for stable maintenance of the cell line and production of monoclonal antibodies. Specific techniques for preparing monoclonal antibodies are described in Antibodies: 20 Laboratory Manual, Harlow and Lane, eds., Cold Spring Harbor Laboratory, 1988.

In addition to monoclonal antibodies and polyclonal antibodies (antisera), the detection techniques of the present invention will also be able to use antibody fragments, such as 25 F(ab), Fv, V_L , V_H , and other fragments. In the use of polyclonal antibodies, however, it may be necessary to adsorb the anti-sera against the target epitopes in order to produce a monospecific antibody population. It will also be possible to employ recombinantly produced antibodies (immunoglobulins) and 30 variations thereof as now well described in the patent and scientific literature. See, for example, EPO 8430268.0; EPO 85102665.8; EPO 85305604.2; PCT/GB 85/00392; EPO 85115311.4; PCT/US86/002269; and Japanese application 85239543, the disclosures of which are incorporated herein by reference. It 35 would also be possible to prepare other recombinant proteins which would mimic the binding specificity of antibodies prepared as just described.

VII. KITS

This invention also provides kits for performing assays of the invention. The kits include means for detecting specifically $A\beta(x-\ge 41)$ and means for detecting specifically $A\beta(x-\le 40)$. The means can include any means known or described above, e.g., binding substances.

In one embodiment, the kit includes a binding substance specific for $A\beta(x-\ge41)$ (i.e., that does not cross react with $A\beta(\le40)$), and a binding substance specific for 10 $A\beta(\le40)$ (i.e., that does not cross react with $A\beta(x-\ge41)$). Such kits are useful, e.g., in pulse-chase assays.

In another embodiment, the kit is useful for immunoassays including two antibodies for each antigen. For example, the kit can further comprise a binding substance specific for the junction region of $A\beta$. Such antibodies are useful for the capture or detection of both $A\beta(x-\ge 41)$ and $A\beta(\le 40)$. In one kit useful for a sandwich ELISA, the binding substance specific for the junction region is bound to a solid phase, and the binding substances specific for $A\beta(x-\ge 41)$ and $A\beta(\le 40)$ are detectably labeled.

The detectable labels can be any known and used in the art including, e.g., a biotinylation label, a radioactive label, a light scattering label, an enzymatic label, a fluorescent label and the like. When the label is enzymatic, the kit can further comprise a substrate for the enzyme.

VIII. <u>TEST COMPOUNDS</u>

The test compounds can be any molecule, compound, or other substance which can be added to the cell culture or administered to the test animal without substantially interfering with cell or animal viability. Suitable test compounds may be small molecules (i.e., molecules whose molecular mass is no more than 1000 Daltons), biological polymers, such as polypeptides, polysaccharides, polynucleotides, and the like. The test compounds will typically be administered to the culture medium at a concentration in the range from about 1 nM to 1 mM, usually from about 10 µM to 1 mM. The test compounds will typically be

administered at a dosage of from 1 ng/kg to 100 mg/kg, usually from 10 μ g/kg to 1 mg/kg.

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Test compounds which are able to inhibit secretion or production of $A\beta(x-\ge41)$ are considered as candidates for further determinations of the ability to block β -amyloid production in animals and humans. Such compounds can be tested in *in vivo* studies, as described below. Inhibition of secretion or production indicates that cleavage of $A\beta$ between amino acids 42/43 has likely been at least partly blocked, reducing the amount of $A\beta(x-\ge41)$ available for forming β -amyloid plagues.

IX. <u>IN VIVO SCREENING</u>

Animal models currently are being used to study Alzheimer's disease. (See, e.g., International Patent Application WO 93/14200, U.S. patent application 08/143,697, filed October 27, 1993 (Atty. docket 4.20, allowed), U.S. Patent 5,387,742, and U.S. application 08/486,538, filed June 7, 1995.) These models are useful for screening compounds that alter the production of $A\beta(x-\ge41)$ in the assays of this invention for their ability to affect the course of Alzheimer's disease, both to ameliorate and aggravate the condition. Transgenic mammalian models, more particularly, rodent models and in particular murine, hamster and guinea pig models, are suitable for this use.

A preferred non-human transgenic animal is one whose cells harbor a PDAPP construct. A PDAPP construct is a nucleic acid construct that comprises a mammalian promoter operatively linked to a cDNA-genomic DNA hybrid coding for the expression of APP. The cDNA-genomic DNA hybrid contains a cDNA sequence encoding APP770 or a cDNA sequence encoding APP770 with a naturally occurring mutation (e.g., a Hardy mutation or the Swedish mutation) substituted with genomic DNA sequences. The genomic DNA sequences consist of exon 6 and an amount of the adjacent downstream intron sufficient for splicing, the KI and OX-2 coding region and an amount of each of their upstream and downstream introns sufficient for splicing, and exon 9 and an amount of the adjacent upstream intron sufficient for splicing,

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substituted into the corresponding region of the cDNA sequence encoding APP770, or the cDNA encoding APP770 with a naturally occurring mutation. The construct is transcribed and differentially spliced in mammalian cells to form mRNA molecules that encode and that are translated into APP695, APP751 and APP770. In certain embodiments, the construct contains a PDGF-β promoter operatively linked with a hybrid sequence encoding an APP gene harboring a Hardy mutation (V717F), and the SV40 polyadenylation signal. One version of the PDAPP construct is presented in Example IX.

Another useful non-human animal model harbors a copy of an expressible transgene sequence which encodes the Swedish mutation of APP (asparagine 595 -leucine 596). The sequence generally is expressed in cells which normally express the naturally-occurring endogenous APP gene (if present). Such transgenes typically comprise a Swedish mutation APP expression cassette, in which a linked promoter and, preferably, an enhancer drive expression of structural sequences encoding a heterologous APP polypeptide comprising the Swedish mutation.

for example, brain homogenate.

The transgenic animals that are usually produced by introducing the transgene or targeting construct into a fertilized egg or embryonic stem (ES) cell, typically by
25 microinjection, electroporation, lipofection, or biolistics.
The transgenic animals express the Swedish mutation APP gene of the transgene (or homologously recombined targeting construct), typically in brain tissue. Preferably, one or both endogenous APP alleles is inactivated and incapable of expressing the
30 wild-type APP.

In all cases, it will be necessary to obtain a control value which is characteristic of the level of $A\beta(x-\ge 41)$ and/or total $A\beta$ or $A\beta(x-\le 40)$ production in the test animal in the absence of test compound(s). In cases where the animal is sacrificed, it will be necessary to base the value on an average or a typical value from other test animals which have been transgenically modified to express the Swedish mutant of human APP but which have not received the administration of any

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test compounds or any other substances expected to affect the level of production of $A\beta(x-\ge41)$ and/or total $A\beta$ or $A\beta(x-\le40)$. Once such control level is determined, test compounds can be administered to additional test animals, in which deviation 5 from the average control value would indicate that the test compound had an effect on the γ -secretase activity in the animal. Test substances which are considered positive, i.e., likely to be beneficial in the treatment of Alzheimer's disease or other β -amyloid-related conditions, will be those which are 10 able to reduce the level of $A\beta(x-241)$ production, preferably by at least 20%, more preferably by at least 50%, and most preferably by at least 80%.

The following examples are offered by way of 15 illustration, not by way of limitation.

EXPERIMENTAL

I. ANTIBODY PREPARATION

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Monoclonal Antibodies to the $A\beta$ Junction Region Monoclonal antibodies to the junction region of $\mathtt{A}eta$ were prepared using a synthetic peptide spanning amino acid residues 13-30, except that AI, amino acids 30 and 31, were substituted with GC (the "junction peptide"). The junction peptide was conjugated to an immunogen (lpha-CD3 ϵ antibody; Clone 25 No. 145-2C11, Boehringer-Mannheim) using m-maleimidobenzoyl-Nhydroxysuccinimide ester (MHS) according to the manufacturer's (Pierce) instructions.

A/J mice were immunized initially intraperitoneally (IP) with the $A\beta$ conjugate mixed with complete Freund's 30 adjuvant. Fourteen days later, the mice were boosted IP with the Aeta conjugate mixed with phosphate buffered saline (PBS) at 14 day intervals. After six total boosts, the mice were finally boosted intravenously with $A\beta$ conjugate mixed with incomplete Freund's adjuvant and fused 3 days later. Fusion of spleen cells with P3.653 myeloma cells was performed as described in Oi and Herzenberg, Selective Methods in Cellular Immunology, Mishell and Shigii, Eds., W.H. Freeman and Company, San Francisco, Chapter 17 (1980). Serum titers and initial

screens were performed by the RIA method described below. Several clones were expanded to a 24-well plate and subjected to further analysis as described below. Clones of interest were produced in mouse ascites.

The RIA method used to screen serum bleeds and fusion hybridoma supernatants was based upon a method developed by Wang et al. (1977) J. Immunol. Methods 18:157-164. Briefly, the supernatant (or serum) was incubated overnight at room temperature on a rotator with $^{125}\text{I-labeled}~\text{A}\beta_{\text{1-28}}$ and Sepharose® 10 4B beads to which sheep anti-mouse IgG had been coupled via cyanogen bromide. The beads from each well were harvested onto glass fiber filter discs with a cell harvester and washed several times with PBS. The filter discs were then transferred to gamma tubes and the bound radioactivity was counted in a 15 gamma counter.

All hybridomas were tested for binding to $A\beta_{1-28}$ using the method described above in the initial screen, and then retested 3 days later. A β_{1-28} positive clones were further characterized for reactivity to ^{125}I -labeled $A\beta_{1-16}$ using 20 the RIA method described above. No clones were found to bind $A\beta_{1-16}$. In a peptide capture ELISA, all clones were found to react with $A\beta_{13-28}$ while no clones reacted with $A\beta_{17-28}$. Therefore, it was determined that all clones had an epitope within the junction region spanning amino acids 16 and 17.

Based on results of the above assays, several clones were expanded into 24 well plates. These clones were further characterized by saturation analysis. Supernatants at the 50% titer point (as determined by the RIA method described above) were added to wells containing Sepharose®-sheep anti-mouse IgG 30 beads, a constant amount of ^{125}I -labeled $A\beta_{1-28}$, and varying amounts of unlabeled junction peptide or $A\beta_{17-28}$. concentration of cold peptide for 50% inhibition was determined for each antibody. For the $A\beta_{17-28}$, no inhibition was seen at 100 ng/well for any clones. The 50% inhibition point for 35 junction peptide ranged from 10-80 ng/well. The clones were also characterized based on reactivity in Western blots. Based on titer point, sensitivity (as determined by the 50% inhibition point), and reactivity on Western blot, several

clones were produced in ascites. Antibodies from hybridoma designated 266 (the "266 antibody") was selected for use as a capture antibody in the assays described below.

Supernatants from wells containing hybridoma cells were screened for antibody with the ability to capture ^{125}I labeled A β 1-42 in solution by immunoprecipitation (Notebook 1101).

B. 2G3 Production

10 Antibody 2G3, specific for $A\beta(x-40)$, was produced by injecting female A/J mice intraperitoneally with 100 μ g immunogen per injection. The immunogen consisted of the peptide NH_2 -Cys-NH- CH_2 -(CH_2)₅-CO-GLMVGGVV-COOH (SEQ ID NO:5), coupled to sheep anti-mouse IgG using maleimidohexanoyl-N-15 hydroxysuccinimide. The immunogen was emulsified with Freund's complete adjuvant for the first immunization, and all subsequent immunizations were with 100 μ g of immunogen emulsified with Freund's incomplete adjuvant at approximately two week intervals.

Three days before fusion, a mouse was boosted with PBS solutions containing 50 μ g immunogen intravenously and 50 μ g intraperitoneally in PBS. The mouse was sacrificed, the spleen was removed, splenocytes were isolated and fused with the SP2/0 mouse myeloma using a modification of the method of Koehler and Milstein.

Supernatants from wells containing hybridoma cells were screened for the ability to produce antibody which recognizes $A\beta(1-40)$ which had been coated onto an ELISA plate. "Positives" were further screened for their ability to capture 30 ^{125}I $A\beta(1-40)$ in solution by immunoprecipitation.

C. Production of 21F12

Antibody 21F12, specific for Aβ(x≥41) was produced by immunizing A/J mice intraperitoneally with 100 μg of immunogen per injection. The immunogen consists of the synthetic peptide NH₂-Cys-NH-CH₂(CH₂)₅-CO-GLMVGGVVIA-COOH (SEQ ID NO:4) coupled to sheep anti-mouse IgG using maleimidohexanoyl-N-hydroxysuccinimide (MHS). The immunogen was emulsified with

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Freund's complete adjuvant for the first immunization, and all subsequent immunizations were with 100 μg of immunogen emulsified with Freund's incomplete adjuvant at approximately two week intervals.

Three days before the fusion, a mouse was injected 5 with 50 μg of immunogen each intravenously and intraperitoneally of immunogen in PBS. Three days post injection, the spleen was removed, splenocytes were isolated and fused with SP2/0 following a modification of the method of 10 Koehler and Milstein.

Specificity of 2G3 and 21F12 D.

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The specificities of antibody 2G3 and 21F12 are demonstrated in Figures 1 and 2. In this assay, antibody 2G3 15 or 21F12 was coated into the wells of an ELISA plate by diluting the purified antibody to a concentration of 10 $\mu \mathrm{g/ml}$ in Well-coating Buffer (0.01 M PO $_4$ pH 8.5), and pipetting 100 μ l of the antibody solution into each well. The solution was left overnight at room temperature and then was removed by 20 aspiration. The non-specific sites of the well were blocked by the addition of 200 μ l 0.25% Human Serum albumin in PBS and incubated for at least one hour at room temperature. blocking solution was removed and the wells were washed one time with wash buffer (Tris buffered Saline, 0.05% Tween 20).

Standards containing between 80-20,000 pg/ml of either $A\beta(1-40)$ or $A\beta(1-42)$ were then prepared by dilution in Specimen Diluent (1 mM PO4, 0.15 M NaCl, pH 7.4, 0.6% Bovine Serum albumin, globulin-free, 0.05% Triton X-405 and 0.05% Thimerosal), and 100 μl of each of these standards were added 30 to the appropriate wells. The standards were incubated for one hour at room temperature, then aspirated and the wells washed four times with wash buffer.

100 μ l of a second antibody (the reporter antibody) was added at a concentration of 0.5 $\mu g/ml$ in specimen diluent. 35 This reporter antibody is biotinylated 3D6 (which recognizes $A\beta$ (1-5)) prepared by the reaction of antibody with NHS-biotin This was allowed to incubate one hour at room temperature, and then washed four times with wash buffer.

To each well, 100 μ l of a 1/5000 dilution of avidin HRP (Vector Labs) was added and allowed to incubate one hour at room temperature. The wells were washed four times in wash buffer, and 100 μ l of Slow TMB (Pierce) were added to each well and incubated fifteen minutes. The reaction was stopped by the addition of 25 μ l of 2 M $\rm H_2SO_4$ and the plates were read at 450-650 on a Vmax reader (Molecular Devices) (Notebook 1344).

As can be seen in Figure 1, antibody 2G3 reacts strongly with $A\beta(1-40)$, but has essentially no cross-reactivity with $A\beta(1-42)$. In Figure 2, it is shown that antibody 21F12 similarly has very high specificity, in this case for $A\beta(1-42)$ over $A\beta(1-40)$. At a concentration of 20,000 pg/ml less than 0.4% of cross reactivity is observed.

15 II. <u>ELISA ASSAY</u>

A. Binding of Capture Antibody to Microtiter Wells

A monoclonal antibody against Aβ(x≥41) or Aβ(x≤40) is diluted to a concentration of 10 μg/ml in a buffer containing 0.23g/L NaH₂PO₄·H₂O, 26.2g/L Na₂HPO₄·7H₂O, 1g/L NaN₃, pH 8.5. One hundred μl/well of this solution is then dispensed in a 96 well white Dynatech Microlite 2, 96 well flat-bottomed plate. The plates are sealed and incubated overnight at room temperature. Following coating, the remaining solution was aspirated and the non-specific binding sites are blocked with 200 μL per well of (NaH₂PO₄·H₂O) 0.2g/L, Na₂HPO₄·7H₂O 0.8g/L, human serum albumin (HSA) crystallized and lyophilized 2.5g/L, pH 7.4. These plates are blocked by incubating for 1 hour at room temperature in the blocking solution.

30 B. <u>Assay Protocol</u>

The calibrators are prepared from a stock solution of $A\beta_{1-42}$, $1\mu g/ml$, in DMSO. In specimen diluent ((NaH₂PO₄·H₂O) 0.2g/L, Na₂HPO₄·7H₂O 2.16g/L, NaN₃ 0.5g/L, bovine serum albumin (BSA) (globulin free) 6g/L, triton x-405 0.5ml/L NaCl 8.5g/L, pH 7.4.), the highest calibrator, 1000 pg/ml (10 μ l $A\beta_{1-42}$ stock (1 μ g/ml DMSO) in 10 ml casein specimen diluent) is prepared. Sequential dilutions are made in specimen diluent to obtain 500, 250, 125, 62.5 and 31.25 pg/ml concentrations of $A\beta_{1-42}$.

One hundred μL per well calibrators or samples are applied to the microtiter plates. The plates are sealed and incubated for 1 hour at room temperature. The plates are then washed three times with washing buffer (NaCl 80 g/L, KCl 3.85 g/L, Tris-HCl 31.75 g/L, tween-20 0.5 ml/L, pH 7.5).

Antibody is diluted in specimen diluent to $1\mu g/ml$ and $100\mu l$ is added per well. The plate is covered and incubated for 1 hour at room temperature. The plate is washed three times with washing buffer. The alkaline phosphatase affinity 10 purified F(ab')2 fragment donkey anti-rabbit IgG (H+L) (Jackson) is diluted 1:1000 in specimen diluent. One hundred μ l/well is added. The plate is covered and incubated for 1 hour at room temperature. The plate is washed three times with washing buffer, then 100µl/well of chemiluminescent 15 substrate is added. The chemiluminescent substrate is prepared by diluting the chemiluminescent reagent, AMPPD (Tropix), and an enhancer, emerald green (Tropix), 1:1000 and 1:100 respectively in 1M diethanolemine buffer, pH 10, containing 1 mM MgCl, and 0.2% NaN_3 . The plates are sealed and incubated for 20 10 to 15 minutes at room temperature. Solution is not aspirated. This time may have to be optimized for different antibody lots.

Chemiluminescence is read and expressed as relative chemiluminescence units (CLU) after 15 minutes using a Dynatech 25 ML 1000.

III. INHIBITION OF APP PROCESSING BY THE MDL 28170 INHIBITOR

We first set out to reproduce the published results of Higaki et al. (1995) Neuron, 14:651-659 on the action of the compound MDL 28170 on APP processing, using human kidney 293 cells stably expressing APP₆₉₅ with the Swedish FAD mutation (K695sw cells). Our experiments were done using a pulse-chase paradigm: K695sw cells were labeled for 2 hours with 35s-methionine and then chased for 2 hours in the presence of 200 μM MDL 28170. Aliquots of the chase media from treated and untreated cells were subjected to SDS-PAGE. No significant differences in the amounts of the major secreted cellular proteins were detected (Fig. 3, lanes 1, 2), suggesting that

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under the conditions of the experiment, MDL 28170 does not interfere with general protein secretion.

We next analyzed the chase media for changes in the amounts of α - and β -cleaved APP_s, using antibodies specific for 5 each form. Antibody 1736 specifically immunoprecipitates α -cleaved APP_s (Haass et al. (1994) J. Biol. Chem. 269:17741-17748; Haass et al. (1995) Nature Med. 1:1291-1296). This antibody revealed an increase in α -APP_s production upon treatment (Fig. 3, lanes 3, 4), indicating that MDL 28170 does not significantly inhibit α -secretase.

192sw specifically immunoprecipitates the β -cleaved APP_s species ending with the Swedish mutant met₅₉₆ (Knops et al. (1995) J. Biol. Chem. 270:2419-2422; Haass et al. (1995) Nature Med. 1:1291-1296). Immunoprecipitation with this antibody showed that MDL 28170 does not significantly inhibit β -secretase activity (Fig. 3, lanes 5, 6).

We next analyzed lysates of the K695sw cells for changes in cellular full-length APP and its C-terminal fragments using antibody C7, directed to the last 20 amino acids of APP (Podlisny et al. (1991) Am. J. Pathol. 138:1423-1435). This antibody precipitates N'- and N'/O'- glycosylated full-length APP and its 10 kDa C-terminal fragment (residues 613-695 of APP₆₉₅) that remains membrane bound after α-secretase cleavage. Upon treatment with MDL 28170, a striking increase in the level of the 10 kDa C-terminal fragment was observed (Fig. 3, lanes 7, 8).

In K695sw cells, the 12 kDa C-terminal fragment (residues 597-695 of APP₆₉₅) which remains membrane bound after β -secretase cleavage cannot be easily resolved and detected by antibody C7 (Citron et al., 1995). We therefore precipitated this fragment using antibody 1282 raised to synthetic $A\beta(1-40)$. This antibody can precipitate the 12 kDa C-terminal but not the 10 kDa fragment, and therefore the faint 12 kDa band is not overshadowed by the much more abundant 10 kDa band. Whereas no 12 kDa fragment was detectable in untreated cells, this band was clearly observed upon treatment with MDL 28170 (Fig. 3, lanes 9, 10).

In summary, no significant inhibition of α - or β -secretase cleavage, but an increase in both the 10 and 12 kDa C-terminal fragments was observed upon treatment with MDL 28170, strongly supporting the role of this compound as a γ -secretase inhibitor which inhibits the conversion of the 10 and 12 kDa fragments to the p3 and A β peptides, respectively.

IV. MDL 28170 INHIBITS THE PRODUCTION OF $A\beta(40)$ AND p3(40) BUT NOT $A\beta(42)$ AND p3(42)

MDL 28170 had previously been shown to inhibit the secretion of $A\beta$ and p3 and therefore had been suggested to inhibit γ -secretase. This inhibition was observed by immunoprecipitating media from treated cells with a polyclonal antibody raised to synthetic $A\beta(1-40)$ (Higaki et al., (1995) Neuron, 14:651-659). Since the vast majority of secreted $A\beta$ and p3 peptides end at amino acid 40, this experiment does not distinguish whether only the major γ -secretase cleavage at position 40 is inhibited.

To address this question, we performed pulse-chase 20 experiments on K695sw cells using different doses of MDL 28170, followed by sequential immunoprecipitation of the same media first with 21F12, a monoclonal antibody raised against amino acids 33-42 of $A\beta$ (described above) which specifically precipitates Aeta peptides ending at position 42, and then with 25 antibody 1282 which precipitates all forms of $A\beta$ and p3 (Haass et al., (1992b) Nature 359:322-325) (Fig. 4A). Interestingly, 21F12 precipitated not only $A\beta$, but also p3 peptides, thus demonstrating the existence of secreted p3(42), which had not been described before. Total $A\beta$ and total p3 were strongly and 30 significantly decreased with doses of MDL 28170 > 50 μ M (e.g., at 200 μM p<0.001). In contrast, $A\beta$ (42) and p3(42) showed a bell-shaped dose response curve, with only a small and insignificant decrease at 200 μM , the dose used by Higaki et al. (1995) Neuron, 14:651-659, and the highest dose tested 35 here. Using MDL 28170 at 200 μM , the experiment was repeated four times and the results were quantitated by phosphorimaging (Fig. 4B). These data indicate that under the conditions

described above the differential effect is significant for both ${\rm A}\beta$ and p3.

V. <u>DIFFERENTIAL INHIBITION IS ACCOMPLISHED UNDER A NUMBER OF</u> CONDITIONS

To make sure that the differential effect observed in $A\beta(40)/A\beta(42)$ and p3(40)/p3(42) precipitations is meaningful, we performed a number of control experiments using the K695sw cells. First we treated K695sw cells in a 2 h pulse 2 h chase 10 paradigm with 1 μM of the phorbolester PDBu, which has been shown to decrease total $A\beta$ but increase total p3, probably by diverting β APP substrate from the β -secretase pathway to the α secretase route (Buxbaum et al., (1993) "Protein phosphorylation inhibits production of alzheimer amyloid $\beta/A4$ 15 peptide, " Proc. Natl. Acad. Sci. USA 90:9195-9198; Hung et al., (1993) "Activation of protein kinase C inhibits cellular production of the amyloid β -protein, " J. Biol. Chem. 268:22959-22962). This effect should be independent of the subsequent γ secretase cleavage and thus the 40 and 42 forms of each 20 metabolite should be equally decreased or increased if the immunoprecipitation paradigm used here works correctly. Indeed, when conditioned media of PDBu-treated cells was precipitated with 21F12, the expected decrease in $A\beta(42)$ and increase in p3(42) were observed indicating that $A\beta(42)$ 25 immunoprecipitation signal does reflect changes in the amounts of precipitable material. Subsequent immunoprecipitation with R1282 shows the same effect for total $A\beta$ and total p3 (Fig. 5A).

The statement that $A\beta(42)$ and p3(42) are not decreased by MDL 28170 depends critically on the quality of the 21F12 antibody. To confirm the effects observed with this antibody, two other previously published $A\beta(42)$ -specific antibodies were used in the pulse chase paradigm with MDL 28170 at 200 μ M. The monoclonal antibody BC05 has been extensively used in ELISA assays to detect $A\beta(42)$ (Asami-Odaka et al., (1995) Biochemistry 34:10272-10278; Gravina et al., (1995) J. Biol. Chem. 270:7013-7016; Suzuki et al., (1994) Science 264:1336-1340).

When media from MDL 28170-treated K695sw cells was precipitated with this antibody, we observed an actual increase in both $A\beta$ (42) and p3(42). The subsequent precipitation with R1282 showed the usual decrease in total $A\beta$ and p3 (Fig. 5B).

The polyclonal antibody C42 has also been shown to be specific for $A\beta(42)$ (Saido et al., (1994) Spatial resolution of the primary β -amyloidogenic process induced in postischemic hippocampus. J. Biol. Chem. 269:15253-15257). Likewise, this antibody did not show a decrease in $A\beta(42)$ and $\beta(42)$ upon treatment whereas the subsequent precipitation with R1282 showed the usual decrease in total β and $\beta(42)$.

The decrease in $A\beta(40)$ and p3(40) was also found when the monoclonal antibody, 2G3 (described above) specific for the free carboxyl-terminus of $A\beta(40)$ and p3(40) was used to precipitate first, followed by 21F12 (Fig. 5D). As expected, the differential inhibition of $A\beta$ production by MDL 28170 was also detected when the precipitations were carried out not sequentially (as described above) but in parallel after the standard pulse-chase. That is, aliquots of media from treated cells and from untreated cells were precipitated with 21F12 for the $A\beta42$ forms and other aliquots were precipitated with antibody 1282 for total $A\beta$ and total p3. This parallel precipitation produced the same result as the sequential precipitations described above (Fig. 5E).

In summary, three different $A\beta(42)$ end-specific antibodies show that MDL 28170 does not strongly decrease $A\beta(42)$ and p3(42) production whereas a monoclonal antibody to $A\beta(40)$ and p3(40) and different polyclonal antibodies to total $A\beta$ and p3 show a strong decrease. Finally, immunoprecipitation with 21F12 followed by antibody 1282 again revealed this differential inhibition when the inhibitor (100 μ M) was applied during a 3 h labeling period instead of in a pulse-chase format (Fig. 5F).

35 VI. <u>DIFFERENTIAL INHIBITION IS OBSERVED IN SEVERAL CELL LINES</u>

To check whether the differential inhibition is
specific for K695sw, three additional cell lines were treated
in the standard 2 h pulse-2 h chase paradigm, and the

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conditioned media were precipitated first with 21F12 and then with antibody 1282. The kidney cell line K695,171 expresses $\ensuremath{\mathsf{APP}}_{\ensuremath{\mathsf{695}}}$ carrying the 717I mutation. This line was chosen because it produces increased levels of $A\beta$ (42) due to the mutation 5 (Suzuki et al., (1994) Science 264:1336-1340). At 200 μM MDL 28170, no decrease of $A\beta$ (42) and p3(42) was observed, whereas $A\beta(40)$ and p3(40) were substantially reduced. The Chinese Hamster ovary cell line CHO695 transfected with wild-type βAPP_{695} cDNA was treated with 200 μM MDL 28170 10 and only a slight decrease of $A\beta(42)$ and p3(42) was observed, whereas $A\beta(40)$ and p3(40) were substantially reduced. (Fig. 6B). The human neuroblastoma cell-line SKN695 expressing wild type etaAPP $_{695}$ was treated with 200 μ M MDL 28170 (Fig. 6C). While ${
m A}eta$ (42) and p3(42) were slightly increased, total ${
m A}eta$ and total 15 p3 were strongly decreased. Thus, differential inhibition of $A\beta$ (42) vs $A\beta$ (40) and p3 (42) vs p3 (40) production is not only observed in K695sw but also in a cell line with an Alzheimer's disease linked β APP717 missense mutation, in a hamster cell line and in a human neural cell line expressing wild type β APP. 20

VII. EXPERIMENTAL PROCEDURES

A. <u>Cell lines</u>

All transfected cell lines described here carry derivatives of pCMV695, a plasmid carrying APP₆₉₅ under control of the CMV promoter (Selkoe, (1988) Proc. Natl. Acad. Sci. 85:7341-7345) K695sw are human embryonic kidney 293 cells stably transfected with a construct carrying the AD-linked double ("Swedish") mutation K595N/M596L (Citron et al. (1992) Nature, 360:672-674); K695₇₁₇₁ are 293 cells stably transfected with APP₆₉₅ carrying the mutation V717I (valine to isoleucine at position 717 in the APP770 numbering system). CHO695 are Chinese hamster ovary cells (CHO) stably transfected with pCMV695 (Oltersdorf (1990) J. Biol. Chem. 265:4432-4437) SKN695 are SK-N-SH human neuroblastoma cells stably transfected with pCMV695.

B. Pulse-chase experiments and immunoprecipitations
To analyze the effect of MDL 28170 on the processing
of APP, cells were grown to confluence in two 10 cm dishes,
pulse-labeled with 600 μCi of [35S]-methionine in 4 ml of serum5 free medium for 2 hours and then chased for 2 hours with 4 ml
medium containing 10% fetal bovine serum and the indicated
final concentration of MDL 28170 (initially dissolved at 200 mM
in DMSO). Control dishes were treated with DMSO alone.

Conditioned media and cell lysates were analyzed by immunoprecipitation, as described (Haass, et al. Nature 359:322-325). Polyclonal antibody R1736 to residues 595-611 of APP₆₉₅, was used to precipitate α -APP_s (Haass, (1994) J. Biol. Chem. 269:17741-17748). This antibody recognizes an epitope that is specific for the free COOH-terminus of α -cleaved APP_s.

- Polyclonal antibody R1282 was generated to synthetic $A\beta_{1-40}$ (Haass, et al. (1992) Nature 359:322-325). This antibody precipitates total $A\beta$ and p3 (and small, variable amounts of APP_s) from the media of cultured cells (Haass, et al. (1992) Nature 359:322-325). The monoclonal antibody 2G3 was raised to
- peptide C(Aminoheptanoic acid)GLMVGGVV (SEQ ID NO:5) and specifically precipitates $A\beta(40)$ and p3(40). Twenty μg of this antibody were used to immunoprecipitate the chase media of 2 dishes. The monoclonal antibody 21F12 was raised to peptide C(Aminoheptanoic acid)GLMVGGVVIA (SEQ ID NO:4) and specifically
- precipitates $A\beta(42)$ and p3(42). Twenty μg of this antibody were used to immunoprecipitate the chase media of 2 dishes. The monoclonal antibody BC05 specifically detects $A\beta(42)$ and p3(42) (Suzuki et al., (1994) Science 264:1336-1340). The polyclonal antibody C7 against the last 20 residues of the APP
- cytoplasmic tail (Podlisny, (1991) Am. J. Pathol. 138:1423-1435) precipitates N'- and N' plus O'-glycosylated full-length APP as well as its C-terminal proteolytic fragments. The antibody sw192 (Knops et al. (1995) J. Biol. Chem. 270:2419-2422; Haass et al. (1995) Nature Med. 1:1291-1296) specifically precipitates β -cleaved APPs carrying the Swedish mutation.
 - SDS-PAGE of immunoprecipitates of cell extracts or of A $\dot{\beta}$ from media was carried out on 10-20% Tris-Tricine gels (Novex), whereas APP $_{\rm s}$ precipitates were electrophoresed on 10%

SDS-polyacrylamide Tris Glycine gels. All quantitations were performed with a Phosphorimager 400A using Image-QuaNT software (Molecular Dynamics). It should be noted the pulse/chase-immunoprecipitation method allows one to assess any changes in $A\beta$ and p3 simultaneously in the same assay, with each of these peptides being visualized directly in the electrophoretic gel.

VIII. HUMAN NEURONS

Human neurons were cultured as previously described (Seubert et al. Nature (1992) 359:325-327) except that the cells were seeded into 6-well plates in neuronal medium without fetal bovine serum but supplemented with B27 (Gibco). Cells were cultured for 2-3 weeks in serum free medium prior to use.

PDAPP mouse brain cells from 16 day old fetal

15 cerebral cortex were cultured following the protocol for human
neurons except the cells were seeded into 24-well plate
clusters in neuronal medium with 5% fetal bovine serum (Sigma)
and 5% Chang's supplement (Irvine Scientific). Cells were
cultured for 5-7 days prior to being used in experiments.

The procedure for examining the effects of substances on A\$\beta\$ production is as follows. Fresh medium is added to the culture wells and then collected after ~ 24 (8-30) hrs. This is the "control" sample from each well that the treated sample will be compared to. Fresh medium, containing the substance to be tested is then added and again harvested after a further ~ 24 hr (8-30) incubation. After collection of this "treated" sample, a cytotoxicity assay is performed on the cells. To perform the cytotoxicity assay, cells are incubated in media containing thiazolyl blue (MTT, Sigma) at 1 mg/ml for 15 minutes. The media are then discarded, and the precipitates are analyzed by solubilization in a buffer containing 50% DMF and 20% SDS. The solubilized dye was quantitated on a Molecular Devices Vmax.

Control and treated samples of culture media are assayed for total $A\beta$ using a sandwich ELISA consisting of two monoclonal antibodies. The first antibody 266, specific to amino acids 13-28 of $A\beta$, is used as a capture antibody (Seubert et al., Nature, supra). The second antibody, 3D6 which is

specific to amino acids 1-5 of $A\beta$, was biotinylated and served as a reporter antibody. The 3D6 biotinylation procedure employed the manufacturer's protocol for NHS-biotin (Pierce) labeling of immunoglobulins, except 100 mM sodium bicarbonate, 5 pH 8.5, buffer was used. The 3D6 antibody does not recognize secreted APP or full-length APP but does recognize $A\beta$ species that begin at position 1.

The samples were also assayed for $A\beta(42)$ with an $A\beta(42)$ specific sandwich ELISA that employed the monoclonal antibody 21F12, which was generated against amino acids 33-42 of $A\beta$, as the capture antibody. This antibody is specific for longer forms of $A\beta$ since it does not cross-react with $A\beta(1-40)$ in ELISA or competitive radioimmunoassay (RIA). Biotinylated 3D6 is also the reporter antibody in this assay.

The 266 and 21F12 mAbs were coated at 10 μ g/ml into 96-well immunoassay plates (Costar) overnight at room temperature. The plates were aspirated and blocked with 0.25% human serum albumin PBS buffer for at least 1 hour at room temperature, then stored desiccated at 4°C until use. The 20 samples and standards were added to the plates and incubated at room temperature for 1.5 hours.

The biotinylated 3D6 was diluted to 0.5 μ g/ml, and incubated in the wells for 1 hour at room temperature. plates were washed 3 times with wash buffer (Tris buffered 25 saline, 0.05% Tween 20) between each step of the assay. Streptavidin-alkaline phosphatase (Boehringer Mannheim), diluted 1:1000, was added to the wells for the total $A\beta$ assay, and avidin-HRP (Vector) diluted 1:4000 was added to the wells for the $A\beta$ (42) assay. These conjugates were incubated for 30 1 hour at room temperature. For the total $A\beta$ assay the fluorometric substrate 4-methyl-umbelliferyl phosphate was added to the wells for 30 minutes, then read in a Millipore Cytofluor 2350 fluorometer. The colorimetric substrate, Slow TMB-ELISA (Pierce), was added for the $A\beta$ (42) assay and allowed 35 to react for 15 minutes, after which the enzymatic reaction was stopped with 2N H₂SO₄. The plates were read on a Molecular Devices Vmax.

Percent inhibition for both total Aeta and Aeta(42) is defined as:

(1-((treated/control_t)/(untreated/control_v))) x 100%,

where treated = value from treated cells

control = value from the same treated well for the

24 hr period prior to testing

untreated = value from well which received no test

substance

control = value from the same untreated well for

the 24 hr prior to testing

It is optimal to divide the values of the treated and untreated samples by their respective values for the 24 hrs prior to the 24 hr test period, as the act of changing the media alone can effect a 5-10% reduction in $A\beta$ production (comparing the untreated to control.).

Several compounds were screened for their ability to inhibit both total $A\beta$ and $A\beta$ (42). Results are shown in Figs. 13A-130.

IX. PDAPP CONSTRUCT

is destroyed.

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25 A cDNA/genomic APP construct containing introns 6, 7 and 8 is prepared by combining APP cDNA encoding exons 1-6 and 9-18 with genomic APP sequences encoding introns 6, 7 and 8, and exons 7 and 8 (see Figs. 8A-8B). In order to create a splicing cassette small enough for convenient insertion in a pUC vector, two deletions in intronic sequences are made. A deletion is made in intron 6 from position 143 of intron 6 to the BamHI site located upstream of the beginning of exon 7 (1658 bp before the beginning of exon '7). Another deletion is made in intron 8 from the first BamHI site in intron 8 to a 35 site at 263 bp before the beginning of exon 9. confusion, these truncated forms of APP introns 6 and 8 are referred to herein as intron 46 and 48. BamHI sites are engineered at the sites of these deletions, so that they are marked by the presence of BamHI sites. In this construct. 40 referred to as PDAPP, exons 7 and 8 and intron 7 are intact genomic sequences, except that the unique XhoI site in intron 7

DNA fragments containing the truncated introns are generated as follows: a BamHI site is engineered 143 bp into intron 6 nucleotide by PCR mutagenesis ("Mutagenesis by PCR" in PCR Technology: Current Innovations (Griffith and Griffith, eds., CRC Press, 1994) pages 69-83) and another BamHI site is engineered by PCR mutagenesis 263 bp prior to the beginning of exon 9. These sites are engineered into separate APP genomic DNA clones containing the junctions of exon 6 and intron 6, and intron 8 and exon 9, respectively, resulting in modified APP genomic DNA clones.

The entire cassette is assembled in the APP cDNA clone as follows (Figure 9). The 889 bp BamHI to XcmI fragment of APP cDNA containing exons 1 through 5 and part of exon 6 (including nucleotides 1 to 843 of Fig. 10 (SEQ ID NO:2)) is 15 cloned into a vector containing BamHI and XhoI sites downstream from the insertion site to make APP770x-oligo-x. APP770xoligo-x is then cut with XcmI and BamHI. Then two fragments are obtained from the modified APP genomic DNA clone containing the junction of exon 6 and intron 6 described above by cutting 20 with XcmI and BamHI. The resulting 34 bp fragment from the XcmI in exon 6 to the XcmI in intron 6, and 131 bp fragment from the XcmI in intron 6 to the artificially created BamHI site at position 143 bp of intron 6 are ligated into APP770xoligo-x in a three-way ligation step to make APP 770x-E6oligo-The orientation of the fragments are confirmed by 25 x. sequencing. APP770x-E6oligo-x is then cut with BamHI and XhoI. Then the 313 bp BamHI and XhoI fragment from the modified APP genomic DNA clone containing the junction of intron 8 and exon 9 is ligated into APP770x-E6oligo-x to make APP770xE6E9x.

APP770xE6E9x is then cut with BamHI and the 6.8 kb BamHI fragment of APP genomic DNA encoding the KPI and OX-2 domains (exons 7 and 8) is inserted at this site. This fragment starts at the BamHI site 1658 bp upstream of the start of exon 7 and extends to the first BamHI site in intron 8. This BamHI fragment is obtained from a lambda phage genomic clone encoding this portion of the APP gene, that was obtained

from a Human Placental genomic library in the Lambda FIXII vector obtained from Stratagene. This BamHI fragment

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originally contained an *XhoI* site which was destroyed by cutting fill in and relegation. The locations of the deletions are diagramed in Figure 11. This clone, containing exons 1-8 and part of 9, and introns 6, 7 and 8, is termed the "APP splicing cassette." The APP splicing cassette is cut out with *NruI* and *XhoI* and used to replace the *NruI* to *XhoI* cDNA fragment of APP cDNA bearing a Hardy mutation. This mutant form of APP cDNA is produced by converting the G at nucleotide position 2145 to T by site directed mutagenesis. This changes the encoded amino acid from Val to Phe. The resulting construct is a combination cDNA/genomic APP "minigene."

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Sequencing of the 6.8 kb BamHI fragment containing APP exons 7 and 8 derived from the APP genomic clone used to generate this construct showed that intron 7 is 2.6 kb long, 15 and that the first BamHI site in intron 8, the upstream site of the deletion in intron 8 engineered into the APP minigene construct, is 2329 bp downstream from the end of exon 8. does not coincide with the restriction map of the APP gene published by Yoshikai et al. (1990) and Yoshikai et al. (1991). 20 Comparison of their map to our sequence indicates that Yoshikai et al. switched the order of two EcoRI fragments in their restriction mapping. The 1.60 kb EcoRI fragment containing exon 8 is actually upstream of the 1.48 kb EcoRI fragment and the 1.48 kb EcoRI fragment Yoshikai et al. mapped into intron 7 25 is actually in intron 8. We have confirmed this location for the EcoRI fragment containing exon 8 by sizing of PCR generated fragments from human DNA.

This APP minigene is operatively linked to the PDGF-β promoter to provide expression of the APP cDNA/genomic

30 construct in mammalian cells. The PDGF β-chain 5' flanking sequence is inserted upstream of the NruI site at the beginning of the APP minigene. This fragment includes 1.3 kb upstream of the transcription initiation site, where the PDGF-β promoter resides, and approximately 70 bp of 5' untranslated region,

35 ending at the AurII site (Higgins et al. (1994)). The late SV40 polyadenylation signal, carried on a 240 bp BamHI to BclI fragment, is added downstream of the APP minigene. This construct, combining the PDGF-β promoter, the APP splicing

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cassette, a Hardy mutation, and the SV40 polyadenylation signal is referred to as PDAPP (Figure 12).

The present invention provides a novel screening 5 method for determining whether a compound alters the production of $A\beta(x-\ge 41)$ and/or $A\beta(x-\le 40)$ or total $A\beta$. While specific examples have been provided, the above description is illustrative and not restrictive. Many variations of the invention will become apparent to those of skill in the art upon review of this specification. The scope of the invention should, therefore, be determined not with reference to the above description, but instead should be determined with reference to the appended claims along with their full scope of equivalents.

All publications and patent documents cited in this application are incorporated by reference in their entirety for all purposes to the same extent as if each individual publication or patent document were so individually denoted.

47 SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: CITRON, MARTIN SELKOE, DENNIS J. SEUBERT, PETER A. SCHENK, DALE
- (ii) TITLE OF INVENTION: SCREENING COMPOUNDS FOR THE ABILITY TO ALTER THE PRODUCTION OF AMYLOID-BETA PEPTIDE (X->41)
- (iii) NUMBER OF SEQUENCES: 5
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: TOWNSEND & TOWNSEND & CREW LLP
 - (B) STREET: TWO EMBARCADERO CENTER, 8TH FLOOR
 - (C) CITY: SAN FRANCISCO
 - (D) STATE: CALIFORNIA
 - (E) COUNTRY: U.S.A.
 - (F) ZIP: 94111-3834
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk

 - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: PCT (B) FILING DATE:

 - (C) CLASSIFICATION:
- (vii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: STORELLA ESQ., JOHN R.

 - (B) REGISTRATION NUMBER: 32,944
 (C) REFERENCE/DOCKET NUMBER: 15270-000650PC
 - (viii) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (415) 326-2400 (B) TELEFAX: (415) 576-0300
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 43 amino acids
 - (B) TYPE: amino acid (C) STRANDEDNESS:

 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
 - Asp Ala Glu Phe Arg His Asp Ser Gly Tyr Glu Val His His Gln Lys
 - Leu Val Phe Phe Ala Glu Asp Val Gly Ser Asn Lys Gly Ala Ile Ile 20

Gly Leu Met Val Gly Gly Val Val Ile Ala Thr

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2310 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: double
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..2310
- (D) OTHER INFORMATION: /function= "coding region for APP770"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

ATG Met 1	CTG Leu	CCC Pro	GGT Gly	TTG Leu 5	GCA Ala	CTG Leu	CTC Leu	CTG Leu	CTG Leu 10	GCC Ala	GCC Ala	TGG Trp	ACG Thr	GCT Ala 15	CGG Arg	48
GCG Ala	CTG Leu	GAG Glu	GTA Val 20	CCC Pro	ACT Thr	GAT Asp	GGT Gly	AAT Asn 25	GCT Ala	GGC Gly	CTG Leu	CTG Leu	GCT Ala 30	GAA Glu	CCC Pro	96
CAG Gln	ATT Ile	GCC Ala 35	ATG Met	TTC Phe	TGT Cys	GGC Gly	AGA Arg 40	CTG Leu	AAC Asn	ATG Met	CAC His	ATG Met 45	AAT Asn	GTC Val	CAG Gln	144
AAT Asn	GGG Gly 50	AAG Lys	TGG Trp	GAT Asp	TCA Ser	GAT Asp 55	CCA Pro	TCA Ser	GGG Gly	ACC Thr	AAA Lys 60	ACC Thr	TGC Cys	ATT Ile	GAT Asp	192
ACC Thr 65	AAG Lys	GAA Glu	GGC Gly	ATC Ile	CTG Leu 70	CAG Gln	TAT Tyr	TGC Cys	CAA Gln	GAA Glu 75	GTC Val	TAC Tyr	CCT Pro	GAA Glu	CTG Leu 80	240
CAG Gln	ATC Ile	ACC Thr	AAT Asn	GTG Val 85	GTA Val	GAA Glu	GCC Ala	AAC Asn	CAA Gln 90	CCA Pro	GTG Val	ACC Thr	ATC Ile	CAG Gln 95	AAC Asn	288
TGG Trp	TGC Cys	AAG Lys	CGG Arg 100	GGC Gly	CGC Arg	AAG Lys	CAG Gln	TGC Cys 105	AAG Lys	ACC Thr	CAT His	CCC Pro	CAC His 110	TTT Phe	GTG Val	336
ATT Ile	CCC Pro	TAC Tyr 115	CGC Arg	TGC Cys	TTA Leu	GTT Val	GGT Gly 120	GAG Glu	TTT Phe	GTA Val	AGT Ser	GAT Asp 125	GCC Ala	CTT Leu	CTC Leu	384
GTT Val	CCT Pro 130	GAC Asp	AAG Lys	TGC Cys	AAA Lys	TTC Phe 135	TTA Leu	CAC His	CAG Gln	GAG Glu	AGG Arg 140	ATG Met	GAT Asp	GTT Val	TGC Cys	432
GAA Glu 145	ACT Thr	CAT His	CTT Leu	CAC His	TGG Trp 150	CAC His	ACC Thr	GTC Val	GCC Ala	AAA Lys 155	GAG Glu	ACA Thr	TGC Cys	AGT Ser	GAG Glu 160	480

					CAT His											528
GAC Asp	AAG Lys	TTC Phe	CGA Arg 180	GGG Gly	GTA Val	GAG Glu	TTT Phe	GTG Val 185	TGT Cys	TGC Cys	CCA Pro	CTG Leu	GCT Ala 190	GAA Glu	GAA Glu	576
					TCT Ser											624
					GAC Asp											672
GTA Val 225	GTA Val	GAA Glu	GTA Val	GCA Ala	GAG Glu 230	GAG Glu	GAA Glu	GAA Glu	GTG Val	GCT Ala 235	GAG Glu	GTG Val	GAA Glu	GAA Glu	GAA Glu 240	720
GAA Glu	GCC Ala	GAT Asp	GAT Asp	GAC Asp 245	GAG Glu	GAC Asp	GAT Asp	GAG Glu	GAT Asp 250	GGT Gly	GAT Asp	GAG Glu	GTA Val	GAG Glu 255	GAA Glu	768
GAG Glu	GCT Ala	GAG Glu	GAA Glu 260	CCC Pro	TAC Tyr	GAA Glu	GAA Glu	GCC Ala 265	ACA Thr	GAG Glu	AGA Arg	ACC Thr	ACC Thr 270	AGC Ser	ATT Ile	816
GCC Ala	ACC Thr	ACC Thr 275	ACC Thr	ACC Thr	ACC Thr	ACC Thr	ACA Thr 280	GAG Glu	TCT Ser	GTG Val	GAA Glu	GAG Glu 285	GTG Val	GTT Val	CGA Arg	864
GAG Glu	GTG Val 290	TGC Cys	TCT Ser	GAA Glu	CAA Gln	GCC Ala 295	GAG Glu	ACG Thr	GGG Gly	CCG Pro	TGC Cys 300	CGA Arg	GCA Ala	ATG Met	ATC Ile	912
TCC Ser 305	CGC Arg	TGG Trp	TAC Tyr	TTT Phe	GAT Asp 310	GTG Val	ACT Thr	GAA Glu	GGG Gly	AAG Lys 315	TGT Cys	GCC Ala	CCA Pro	TTC Phe	TTT Phe 320	960
TAC Tyr	GGC Gly	GGA Gly	TGT Cys	GGC Gly 325	GGC Gly	AAC Asn	CGG Arg	AAC Asn	AAC Asn 330	TTT Phe	GAC Asp	ACA Thr	GAA Glu	GAG Glu 335	TAC Tyr	1008
TGC Cys	ATG Met	GCC Ala	GTG Val 340	TGT Cys	GGC Gly	AGC Ser	GCC Ala	ATG Met 345	TCC Ser	CAA Gln	AGT Ser	TTA Leu	CTC Leu 350	AAG Lys	ACT Thr	1056
ACC Thr	CAG Gln	GAA Glu 355	CCT Pro	CTT Leu	GCC Ala	CGA Arg	GAT Asp 360	CCT	GTT Val	AAA Lys	CTT Leu	CCT Pro 365	ACA Thr	ACA Thr	GCA Ala	1104
GCC Ala	AGT Ser 370	ACC Thr	CCT Pro	GAT Asp	GCC Ala	GTT Val 375	GAC Asp	AAG Lys	TAT	CTC Leu	GAG Glu 380	ACA Thr	CCT Pro	GGG Gly	GAT Asp	1152
GAG Glu 385	Asn	GAA Glu	CAT His	GCC Ala	CAT His 390	TTC Phe	CAG Gln	AAA Lys	GCC Ala	AAA Lys 395	GAG Glu	AGG Arg	CTT Leu	GAG Glu	GCC Ala 400	1200
AAG Lys	CAC His	CGA Arg	GAG Glu	AGA Arg 405	ATG Met	TCC Ser	CAG Gln	GTC Val	ATG Met 410	AGA Arg	GAA Glu	TGG Trp	GAA Glu	GAG Glu 415	GCA Ala	1248
GAA Glu	CGT Arg	CAA Gln	GCA Ala 420	Lys	AAC Asn	TTG Leu	CCT Pro	AAA Lys 425	Ala	GAT Asp	AAG Lys	AAG Lys	GCA Ala 430	GTT Val	ATC Ile	1296

G11	1 1112	435	GII	. GIU	ı Lys	· vai	440	Ser	Leu	ı Glı	ı Glr	1 Glu 445	ı Alá	a Ala	C AAC a Asn	1344
GI	450	GII	GII	ren	ı vaı	455	Inr	His	Met	Ala	Arg 460	Va]	Glı	ı Ala	ATG Met	1392
465	, ASI	Asp	Arg	Arg	470	Leu	Ala	Leu	ı Glu	475	Tyr	Ile	Thr	Ala	CTG Leu 480	1440
GII	HIG	vai	PIO	485	Arg	Pro	Arg	His	Val 490	Phe	Asn	Met	Leu	Lys 495		1488
ıyı	val	Arg	500	GIU	GIN	Lys	Asp	Arg 505	Gln	His	Thr	Leu	Lys 510	His	TTC Phe	1536
GAG Glu	CAT His	GTG Val 515	CGC Arg	ATG Met	GTG Val	GAT Asp	CCC Pro 520	AAG Lys	AAA Lys	GCC Ala	GCT Ala	CAG Gln 525	ATC Ile	CGG	TCC Ser	1584
CAG Gln	GTT Val 530	ATG Met	ACA Thr	CAC His	CTC Leu	CGT Arg 535	GTG Val	ATT Ile	TAT Tyr	GAG Glu	CGC Arg 540	ATG Met	AAT Asn	CAG Gln	TCT Ser	1632
CTC Leu 545	TCC Ser	CTG Leu	CTC Leu	TAC Tyr	AAC Asn 550	GTG Val	CCT Pro	GCA Ala	GTG Val	GCC Ala 555	GAG Glu	GAG Glu	ATT Ile	CAG Gln	GAT Asp 560	1680
GAA Glu	GTT Val	GAT Asp	GAG Glu	CTG Leu 565	CTT Leu	CAG Gln	AAA Lys	GAG Glu	CAA Gln 570	AAC Asn	TAT Tyr	TCA Ser	GAT Asp	GAC Asp 575	GTC Val	1728
TTG Leu	GCC Ala	AAC Asn	ATG Met 580	ATT Ile	AGT Ser	GAA Glu	CCA Pro	AGG Arg 585	ATC Ile	AGT Ser	TAC Tyr	GGA Gly	AAC Asn 590	GAT Asp	GCT Ala	1776
CTC Leu	ATG Met	CCA Pro 595	TCT Ser	TTG Leu	ACC Thr	GAA Glu	ACG Thr 600	AAA Lys	ACC Thr	ACC Thr	GTG Val	GAG Glu 605	CTC Leu	CTT Leu	CCC Pro	1824
GTG Val	AAT Asn 610	GGA Gly	GAG Glu	TTC Phe	AGC Ser	CTG Leu 615	GAC Asp	GAT Asp	CTC Leu	CAG Gln	CCG Pro 620	TGG Trp	CAT His	TCT Ser	TTT Phe	1872
GGG Gly 625	GCT Ala	GAC As p	TCT Ser	GTG Val	CCA Pro 630	GCC Ala	AAC Asn	ACA Thr	GAA Glu	AAC Asn 635	GAA Glu	GTT Val	GAG Glu	CCT Pro	GTT Val 640	1920
GAT Asp	GCC Ala	CGC Arg	Pro	GCT Ala 645	GCC Ala	GAC Asp	CGA Arg	GGA Gly	CTG Leu 650	ACC Thr	ACT Thr	CGA Arg	CCA Pro	GGT Gly 655	TCT Ser	1968
GGG Gly	TTG Leu	ACA Thr	AAT Asn 660	ATC Ile	AAG Lys	ACG Thr	GIU	GAG Glu 665	ATC Ile	TCT Ser	GAA Glu	GTG Val	AAG Lys 670	ATG Met	GAT Asp	2016
GCA Ala	GAA Glu	TTC Phe 675	CGA Arg	CAT His	GAC Asp	ser	GGA Gly 680	TAT Tyr	GAA Glu	GTT Val	His	CAT His 685	CAA Gln	AAA Lys	TTG Leu	2064
GTG Val	TTC Phe 690	TTT Phe	GCA Ala	GAA Glu	Asp	GTG Val 695	GGT :	TCA Ser	AAC Asn	AAA Lys	GGT Gly 700	GCA Ala	ATC Ile	ATT Ile	GGA Gly	2112

 ATG Met		 		 	 	 		2160
 ATG Met	 					 		2208
 GTT Val								2256
 CAG Gln	 							2304
 AAC Asn 770								2310

(2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 770 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

 Met
 Leu
 Pro
 Gly
 Leu
 Leu
 Leu
 Leu
 Leu
 Leu
 Ala
 Ala
 Trp
 Thr
 Ala
 Arg

 Ala
 Leu
 Glu
 Val
 Pro
 Thr
 Asp
 Gly
 Asn
 Ala
 Gly
 Leu
 Leu
 Ala
 Glu
 Pro
 330
 Pro
 Pro
 Asp
 Gly
 Asp
 Leu
 Asn
 Met
 His
 Met
 Asn
 Val
 Gln
 Asp
 Leu
 Asn
 Met
 His
 Met
 Asn
 Val
 Gln
 Asn
 Met
 His
 Asn
 Val
 Gln
 Asn
 Met
 His
 Asn
 Val
 Gln
 Asn
 Val
 Thr
 Lys
 Thr
 Cys
 Ile
 Asp
 Pro
 Ser
 Glu
 Asn
 Asp
 Pro
 Glu
 Asp
 Pro
 Ile
 Asp
 Asp
 Ile
 Asp
 Ile
 Asp
 Ile
 Asp
 Ile
 Asp
 Ile

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Ser	Asp	Asn 195	Val	Asp	ser	Ala	Asp 200	Ala	Glu	Glu	Asp	Asp 205	Ser	Asp	Val
Trp	Trp 210	Gly	Gly	Ala	Asp	Thr 215	Asp	Tyr	Ala	Asp	Gly 220	Ser	Glu	Asp	Lys
Val 225	Val	Glu	Val	Ala	Glu 230	Glu	Glu	Glu	Val	Ala 235	Glu	Val	Glu	Glu	Glu 240
Glu	Ala	Asp	Asp	Asp 245	Glu	Asp	Asp	Glu	Asp 250	Gly	Asp	Glu	Val	Glu 255	Glu
Glu	Ala	Glu	Glu 260	Pro	Tyr	Glu	Glu	Ala 265	Thr	Glu	Arg	Thr	Thr 270	Ser	Ile
Ala	Thr	Thr 275	Thr	Thr	Thr	Thr	Thr 280	Glu	Ser	Val	Glu	Glu 285	Val	Val	Arg
Glu	Val 290	Cys	Ser	Glu	Gln	Ala 295	Glu	Thr	Gly	Pro	Cys 300	Arg	Ala	Met	Ile
Ser 305	Arg	Trp	Tyr	Phe	Asp 310	Val	Thr	Glu	Gly	Lys 315	Cys	Ala	Pro	Phe	Phe 320
Tyr	Gly	Gly	Cys	Gly 325	Gly	Asn	Arg	Asn	Asn 330	Phe	Asp	Thr	Glu	Glu 335	Tyr
Cys	Met	Ala	Val 340	Cys	Gly	Ser	Ala	Met 345	Ser	Gln	Ser	Leu	Leu 350	Lys	Thr
Thr	Gln	Glu 355	Pro	Leu	Ala	Arg	Asp 360	Pro	Val	Lys	Leu	Pro 365	Thr	Thr	Ala
Ala	Ser 370	Thr	Pro	Asp	Ala	Val 375	Asp	Lys	Tyr	Leu	Glu 380	Thr	Pro	Gly	Asp
Glu 385	Asn	Glu	His	Ala	His 390	Phe	Gln	Lys	Ala	Lys 395	Glu	Arg	Leu	Glu	Ala 400
Lys	His	Arg	Glu	Arg 405	Met	Ser	Gln	Val	Met 410	Arg	Glu	Trp	Glu	Glu 415	Ala
Glu	Arg	Gln	Ala 420	Lys	Asn	Leu	Pro	Lys 425	Ala	Asp	Lys	Lys	Ala 430	Val	Ile
Gln	His	Phe 435	Gln	Glu	Lys	Val	Glu 440	Ser	Leu	Glu	Gln	Glu 445	Ala	Ala	Asn
Glu	Arg 45 0	Gln	Gln	Leu	Val	Glu 455	Thr	His	Met	Ala	Arg 460	Val	Glu	Ala	Met
Leu 465	Asn	Asp	Arg	Arg	Arg 470	Leu	Ala	Leu	Glu	Asn 475	Tyr	Ile	Thr	Ala	Leu 480
Gln	Ala	Val	Pro	Pro 485	Arg	Pro	Arg	His	Val 490	Phe	Asn	Met	Leu	Lys 495	Lys
Tyr	Val	Arg	Ala 50 0	Glu	Gln	Lys	Asp	Arg 505	Gln	His	Thr	Leu	Lys 510	His	Phe
Glu	His	Val 515	Arg	Met	Val	Asp	Pro 520	Lys	Lys	Ala	Ala	Gln 525	Ile	Arg	Ser
Gln	Val 530	Met	Thr	His	Leu	Arg 535	Val	Ile	Туr	Glu	Arg 540	Met	Asn	Gln	Ser

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- Leu Ser Leu Leu Tyr Asn Val Pro Ala Val Ala Glu Glu Ile Gln Asp 550 Glu Val Asp Glu Leu Leu Gln Lys Glu Gln Asn Tyr Ser Asp Asp Val Leu Ala Asn Met Ile Ser Glu Pro Arg Ile Ser Tyr Gly Asn Asp Ala Leu Met Pro Ser Leu Thr Glu Thr Lys Thr Thr Val Glu Leu Leu Pro 600 Val Asn Gly Glu Phe Ser Leu Asp Asp Leu Gln Pro Trp His Ser Phe Gly Ala Asp Ser Val Pro Ala Asn Thr Glu Asn Glu Val Glu Pro Val Asp Ala Arg Pro Ala Ala Asp Arg Gly Leu Thr Thr Arg Pro Gly Ser Gly Leu Thr Asn Ile Lys Thr Glu Glu Ile Ser Glu Val Lys Met Asp 660 Ala Glu Phe Arg His Asp Ser Gly Tyr Glu Val His His Gln Lys Leu Val Phe Phe Ala Glu Asp Val Gly Ser Asn Lys Gly Ala Ile Ile Gly Leu Met Val Gly Gly Val Val Ile Ala Thr Val Ile Val Ile Thr Leu Val Met Leu Lys Lys Gln Tyr Thr Ser Ile His His Gly Val Val Glu Val Asp Ala Ala Val Thr Pro Glu Glu Arg His Leu Ser Lys Met Gln Gln Asn Gly Tyr Glu Asn Pro Thr Tyr Lys Phe Phe Glu Gln Met Gln Asn
- (2) INFORMATION FOR SEQ ID NO:4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (ix) FEATURE:

- (A) NAME/KEY: Modified-site
- (B) LOCATION: 2
 (D) OTHER INFORMATION: /note= "Xaa = NH CH-2 (CH-2)-5 - CO in an amine linkage between amino acid Cys in position 1 and amino acid Gly in position 3"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Cys Xaa Gly Leu Met Val Gly Gly Val Val Ile Ala

- (2) INFORMATION FOR SEQ ID NO:5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 amino acids
 (B) TYPE: amino acid

 - (C) STRANDEDNESS:
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (ix) FEATURE:
 - (A) NAME/KEY: Modified-site (B) LOCATION: 2
 - (D) OTHER INFORMATION: /note= "Xaa = NH CH-2 (CH-2)-5 - CO in an amine linkage between amino acid Cys in position 1 and amino acid Gly in position 3"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Cys Xaa Gly Leu Met Val Gly Gly Val Val

WHAT IS CLAIMED IS:

- A method for determining whether a compound 1 1. alters the amount of at least one $A\beta(x-\ge41)$ peptide produced by 2 a cell, comprising the steps of: 3 administering the compound to a culture 4 comprising the cell; 5 measuring the amount of the $A\beta(x-\ge 41)$ peptide, 6 specifically, in a sample from the culture; and 7 determining whether the measured amount is 8 different than the amount expected in a sample from a culture 9 comprising the cell to which no compound has been administered; 10 whereby a difference between the measured amount 11 and the expected amount indicates that the compound alters the 12 amount of an $A\beta(x-241)$ peptide produced by the cell. 13
 - The method of claim 1 wherein the amount of the $A\beta(x-\ge 41)$ peptide is measured by immunoassay.
 - The method of claim 2 wherein the immunoassay is a sandwich immunoassay using a capture binding substance bound to a solid phase and a labeled detection binding substance.
 - 1 4. The method of claim 3 wherein the capture 2 binding substance is specific for $A\beta(x-\ge41)$ peptides.
 - 5. The method of claim 4 wherein the binding substance has the specificity of a binding substance raised against peptide NH₂-Cys-NH-CH₂(CH₂)₅-CO-GLMVGGVVIA-COOH (SEQ ID NO:4).
 - 7 6. The method of claim 4 wherein the labeled 8 detection binding substance is specific for an epitope within 9 the junction region of $A\beta$.
 - 7. The method of claim 6 wherein the labeled detection binding substance has the specificity of a binding substance raised against the junction peptide of $A\beta$.

- 1 8. The method of claim 4 wherein the labeled
- 2 detection binding substance is specific for an $A\beta$ peptide whose
- 3 amino-terminal amino acid is amino acid no. 1 of $A\beta$.
- 1 9. The method of claim 3 wherein the capture
- 2 antibody is specific for an epitope within the junction region
- of $A\beta$ and the labeled detection antibody is specific for $A\beta$ (x-
- 4 ≥41) peptides.
- 1 10. The method of claim 3 wherein the culture
- 2 comprises primary human neurons or primary neurons from a
- 3 transgenic mouse harboring a PDAPP construct.
- 1 11. The method of claim 10 wherein the PDAPP
- 2 construct contains the mutation V717F.
- 1 12. The method of claim 2 wherein the step of
- 2 measuring the amount of the $A\beta(x-\ge41)$ peptide, specifically, in
- 3 a sample from the culture comprises:
- 4 pulsing the culture with a radioactive label for
- 5 protein;
- chasing the culture without a radioactive label;
- 7 administering the compound to the cell during
- 8 the chase period;
- 9 contacting a sample from the culture with a
- binding substance specific for $A\beta(x-\ge41)$ peptides; and
- determining the amount of radioactive label
- 12 attached to the binding substance.
 - 1 13. The method of claim 1 wherein the culture
 - 2 comprises primary human neurons or primary neurons from a
 - 3 transgenic mouse harboring the PDAPP construct.
 - 1 14. The method of claim 13 wherein the PDAPP
 - 2 construct contains the mutation V717F.
 - 3 15. The method of claim 1 wherein the culture
 - 4 comprises a 293 human kidney cell line, a human neuroglioma

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- cell line, a human HeLa cell line, a primary endothelial cell 5
- line, a primary human fibroblast line, a primary lymphoblast 6
- line, human mixed brain cells, or a Chinese hamster ovary (CHO)
- cell line. 8
- The method of claim 1 wherein the cell is a host 1
- cell transfected with a recombinant expression vector encoding 2
- 3 a human APP.
- The method of claim 16 wherein the human APP 1 17.
- 2 possesses a Hardy mutation.
- 1 18. The method of claim 1 wherein the cell
- 2 overproduces $A\beta(x-\ge41)$ peptides.
- 1 The method of claim 1 wherein the sample is 19.
- medium conditioned by the cell in culture. 2
- 1 The method of claim 1 further comprising the 20.
- step of determining whether the compound is toxic to the cell. 2
- 1 A method for determining whether a compound 21.
- alters the amount of at least one $A\beta(x-\ge41)$ peptide produced by 2
- a cell and alters the amount of either total ${\tt A}{\beta}$ or at least one 3
- $A\beta(x-\leq 40)$ peptide produced by the cell comprising: 4
- 5 administering the compound to a culture
- 6 comprising the cell;
- measuring the amount of the $A\beta(x-\ge41)$ peptide, 7
- 8 specifically, in a sample from the culture;
- measuring the amount of total $A\beta$ or the $A\beta$ (x-9
- ≤40) peptide, specifically, in a sample from the culture; and 10
- determining whether the measured amounts are 11
- different than the amounts expected in a sample from a culture 12
- comprising the cell to which no compound has been administered; 13
- whereby differences between the measured amounts 14
- and the expected amounts indicate that the compound alters the 15
- amount of the $A\beta(x-241)$ peptide by a cell and/or the amount of 16
- 17 total $A\beta$ or the $A\beta(x-\leq 40)$ peptide by the cell.

- 1 22. The method of claim 21 wherein the amount of the
- 2 $A\beta(x-\ge41)$ peptide and the amount of total $A\beta$ or the $A\beta(x-\le40)$
- 3 peptide are measured by immunoassay.
- 1 23. The method of claim 22 wherein the immunoassay
- 2 is a sandwich immunoassay comprising capture binding substances
- 3 bound to a solid phase and labeled detection binding
- 4 substances.
- 1 24. The method of claim 23 wherein the capture
- 2 binding substance for measuring the amount of the $A\beta(x-\ge 41)$
- 3 peptides are specific for $A\beta(x-241)$ peptides.
- 1 25. The method of claim 24 wherein the binding
- 2 substance specific for $A\beta(x-\ge41)$ peptides has the specificity
- 3 of a binding substance raised against peptide NH₂-Cys-NH-CH₂-
- 4 (CH₂)₅-CO-GLMVGGVVIA-COOH (SEQ ID NO:4).
- 1 26. The method of claim 24 wherein the labeled
- 2 detection binding substance for measuring the amount of $A\beta$ (x-
- 3 ≥41) peptides is specific for an epitope within the junction
- 4 region of $A\beta$ or is specific for an $A\beta$ peptide whose amino-
- 5 terminal amino acid is amino acid no. 1 of $A\beta$.
- 1 27. The method of claim 23 wherein the capture
- 2 binding substance for measuring the amount of total $A\beta$ is
- 3 specific for an epitope within the junction region of $A\beta$ and
- 4 the capture binding substance for measuring the amount of the
- 5 A β (x- \leq 40) peptides is specific for A β (x- \leq 40) peptides.
- 1 28. The method of claim 27 wherein the binding
- 2 substance specific for an epitope within the junction region of
- 3 A β has the specificity of a binding substance raised against
- 4 the junction peptide of $A\beta$, and the binding substance specific
- 5 for $A\beta(x-\leq 40)$ peptides has the specificity of a binding
- 6 substance raised against the peptide NH₂-Cys-NH-CH₂-(CH₂)_s-CO-
- 7 GLMVGGVV-COOH (SEQ ID NO:5).

- 1 29. The method of claim 27 wherein the labeled 2 detection binding substance for measuring the amount of $A\beta$ (x-
- 3 <40) peptides has the specificity of a binding substance raised
- 4 against the junction peptide, and the binding substance for
- 5 measuring the amount of total $A\beta$ has the specificity of a
- 6 binding substance raised against a peptide whose amino acid
- 7 sequence is amino acids within 1-5 of $A\beta$ or amino acids 17-24
- 8 of $A\beta$.
- 1 30. The method of claim 23 wherein the capture
- 2 binding substance for measuring the amount of at least one of
- 3 $A\beta(x-\ge41)$ peptide, total $A\beta$ or $A\beta(x-\le40)$ peptide is specific
- 4 for an epitope within the junction region of $A\beta$.
- 1 31. The method of claim 23 wherein the culture
- 2 comprises primary human neurons or primary neurons from a
- 3 transgenic mouse harboring the PDAPP construct.
- 1 32. The method of claim 22 wherein the step of
- 2 measuring the amount of the $A\beta(x-\ge41)$ peptide, total $A\beta$ or the
 - $A\beta(x-\leq 40)$ peptide in a sample from the culture comprises,
- 4 before administration of the compound:
- 5 pulsing the culture with a radioactive label for
- 6 protein;

- 7 chasing the culture without a radioactive label;
- administering the compound to the cell during
- 9 the chase period;
- 10 contacting a sample from the culture with a
- binding substance specific for at least one $A\beta(x-241)$ peptide;
- contacting a sample from the culture with a
- binding substance specific for total A β or at least one A β (x-
- 14 ≤40) peptide; and
- determining the amount of radioactive label
- 16 attached to the binding substances.
 - 1 33. The method of claim 23 wherein the culture
 - 2 comprises primary human neurons or primary neurons from a
 - 3 transgenic mouse harboring the PDAPP construct.

- 1 34. The method of claim 22 wherein the culture
- 2 comprises a 293 human kidney cell line, a human neuroglioma
- 3 cell line, a human HeLa cell line, a primary endothelial cell
- 4 line, a primary human fibroblast line, a primary lymphoblast
- 5 line, human mixed brain cells, or a Chinese hamster ovary (CHO)
- 6 cell line.
- 1 35. The method of claim 22 wherein the cell is a
- 2 host cell transfected with a recombinant expression vector
- 3 encoding a human APP.
- 1 36. The method of claim 35 wherein the human APP
- 2 carries a Hardy mutation.
- 1 37. The method of claim 22 wherein the cell
- 2 overproduces $A\beta(x-\ge41)$ peptides.
- 1 38. The method of claim 22 wherein the sample is
- 2 medium conditioned by the cell in culture.
- 1 39. The method of claim 22 further comprising the
- 2 step of determining whether the compound is toxic to the cell.
- 1 40. A kit for specifically detecting at least one
- 2 $A\beta(x-41)$ peptide and at least one $A\beta(x-40)$ peptide in a
- 3 sample comprising:
- a binding substance specific for at least one
- 5 $A\beta(x-241)$ peptide; and
- a binding substance specific for at least one
- 7 $A\beta(x-\leq 40)$ peptide.
- 1 41. The kit of claim 40 wherein the binding
- 2 substance specific for $A\beta(x-\ge41)$ peptides has the specificity
- of a binding substance raised against peptide NH2-Cys-NH-CH2-
- 4 (CH₂)₅-CO-GLMVGGVVIA-COOH (SEQ ID NO:4); and the binding
- 5 substance specific for $A\beta(x-\le 40)$ peptides has the specificity
- of a binding substance raised against the peptide NH2-Cys-NH-
- 7 $CH_2-(CH_2)_5-CO-GLMVGGVV-COOH$ (SEQ ID NO:5).

- 1 42. A kit for specifically detecting at least one 2 $A\beta(x-241)$ peptide and either total $A\beta$ or at least one $A\beta(x-540)$
- peptide in a sample in a sandwich immunoassay comprising:
- a) at least two different binding substances for
- 5 measuring the amount of $A\beta(x-\ge41)$ peptide; and
- 6 b) at least two different binding substances for
- 7 measuring the amount of total $A\beta$ or $A\beta(x-\le 40)$ peptides.
- 1 43. The kit of claim 42 wherein one of the binding
- substances for measuring the amount of the $A\beta(x-\ge41)$ peptides
- 3 is a capture binding substance specific for $A\beta(x-\ge41)$ peptides
- 4 bound to a solid phase.
- 1 44. The kit of claim 43 wherein the binding
- substance specific for $A\beta(x-\ge41)$ peptides has the specificity
- of a binding substance raised against peptide $\mathrm{NH_2-Cys-NH-CH_2-}$
- 4 (CH₂)₅-CO-GLMVGGVVIA-COOH (SEQ ID NO:4).
- 1 45. The kit of claim 42 wherein one of the binding
- substances for measuring the amount of $A\beta(x-241)$ peptides is a
- 3 labeled detection binding substance specific for an epitope
- 4 within the junction region of $A\beta$.
- 1 46. The kit of claim 42 wherein one of the binding
- 2 substances for measuring the amount of total $A\beta$ is a capture
- 3 binding substance specific for an epitope within the junction
- 4 region of $A\beta$ bound to a solid phase and one of the binding
- substances for measuring the amount of the $A\beta(x-\le 40)$ peptides
- is a capture binding substance specific for $A\beta(x-\le 40)$ peptides
- 7 bound to a solid phase.
- 1 47. The kit of claim 46 wherein the binding
- 2 substance specific for an epitope within the junction region of
- 3 Aeta has the specificity of a binding substance raised against
- 4 the junction peptide of $A\beta$, and the binding substance specific
- 5 for $A\beta(x-\leq 40)$ peptides has the specificity of a binding

substance raised against the peptide NH₂-Cys-NH-CH₂-(CH₂)₅-CO-GLMVGGVV-COOH (SEQ ID NO:5).

- 48. The kit of claim 45 wherein the labeled detection binding substance for measuring the amount of $A\beta(x-40)$ peptides has the specificity of a binding substance raised against the junction peptide, and the binding substance for measuring the amount of total $A\beta$ has the specificity of a binding substance raised against a peptide whose amino acid sequence is amino acids within 15 of 10 areas in the sequence.
- 7 sequence is amino acids within 1-5 of $A\beta$ or amino acids 17-24 of $A\beta$.
- 1 49. The kit of claim 42 wherein the capture binding 2 substance for measuring the amount of at least one of $A\beta(x-\ge 41)$
- peptide, total A β or A β (x- \leq 40) peptide is specific for an
- 4 epitope within the junction region of $A\beta$.
- 50. The kit of claim 45 wherein the detectable label is a biotinylation label, a radioactive label, a light scattering label, an enzymatic label or a fluorescent label.
- 51. The kit of claim 48 wherein the detectable label is a biotinylation label, a radioactive label, a light scattering label, an enzymatic label or a fluorescent label.
- 52. A method for determining whether a compound alters the amount of at least one $A\beta(x-\ge41)$ peptide produced by a non-human mammal and alters the amount of either total $A\beta$ or at least one $A\beta(x-\le40)$ peptide produced by the non-human mammal comprising:
- measuring a first amount of the $A\beta(x-\ge41)$ peptide in a sample from a non-human animal used as a model of Alzheimer's disease;
- measuring a first amount of total $A\beta$ or the $A\beta(x-\leq 40)$ peptide in a sample from the non-human animal;
- administering the compound to the non-human animal;
- measuring a second amount of the $A\beta(x-241)$ peptide in
- 13 a sample from the non-human animal;

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- measuring a second amount of total $A\beta$ or the $A\beta$ (x-
- 15 <40) peptide in a sample from the non-human animal; and
- 16 comparing the first amounts with the second amounts,
- whereby the comparison indicates whether the compound
- 18 increases, decreases, or leaves unchanged the amount of the
- 19 $A\beta(x-\ge41)$ peptide and increases, decreases, or leaves unchanged
- 20 the amount of the $A\beta(x-\le 40)$ peptide.
- 1 53. The method of claim 52 wherein the non-human
- 2 animal is a rodent.
- 1 54. The method of claim 53 wherein the rodent is a
- 2 mouse.
- 1 55. The method of claim 52 wherein the non-human
- 2 animal harbors a copy of an expressible transgene sequence
- 3 which a Hardy mutation.
- 56. The method of claim 52 wherein the non-human
- 1 animal is transformed to express the Swedish mutation of human
- β -amyloid precursor protein (APP).
- 1 57. The method of claim 52 wherein the non-human
- 2 animal is a transgenic animal harboring a PDAPP construct.
- 1 58. The method of claim 57 wherein the animal is a
- 2 mouse.

STANDARD CURVES OF AB 1-40 AND AB 1-42 WITH 2G3 AS A CAPTURE ANTIBODY

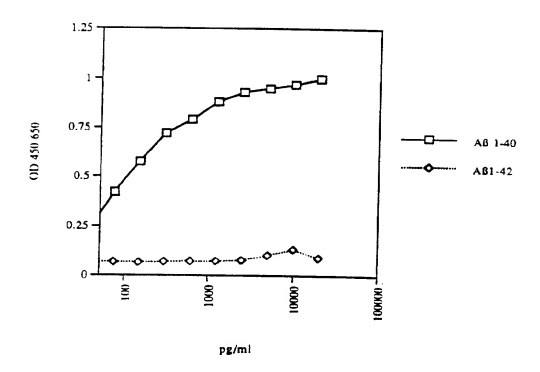


Figure 1

STANDARD CURVES OF AB 1-40 VS AB1-42 USING 21F12 AS A CAPTURE ANTIBODY

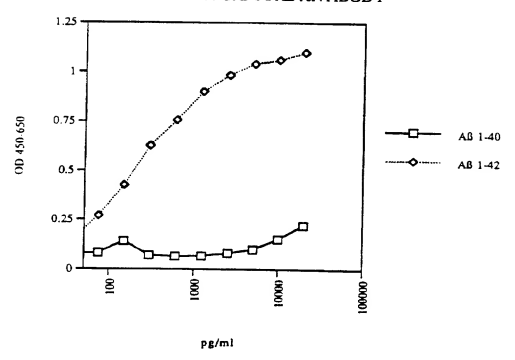


Figure 2

no ab	1736 192sw	C 7	1 28 2
0 200	0 200 0 200 μMMDL	0 200	0 200 μM MDL
		3 = ± N+0	
4: 35:	APP _s		
* *			==
			•••; ••
			← 12 kDa
		7 0	
1 2	3 4 5 6	7 8	9 10

Fig 3

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Α 0 50 100 200 μ**M M**DL

1. 21F12 $-AB_{42}$ $-p3_{42}$

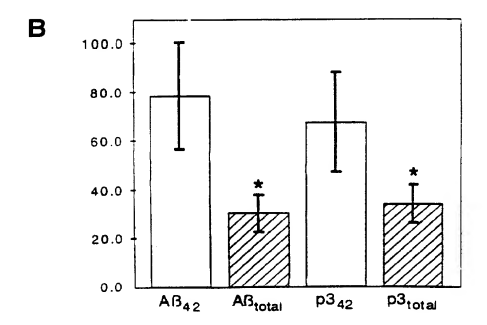


Fig 4

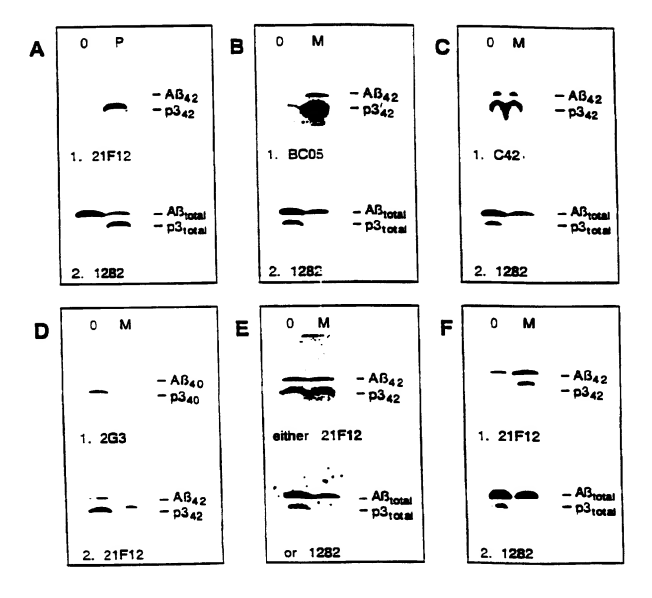


Fig 5

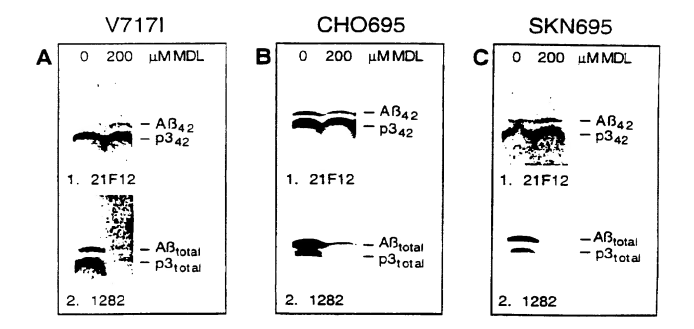
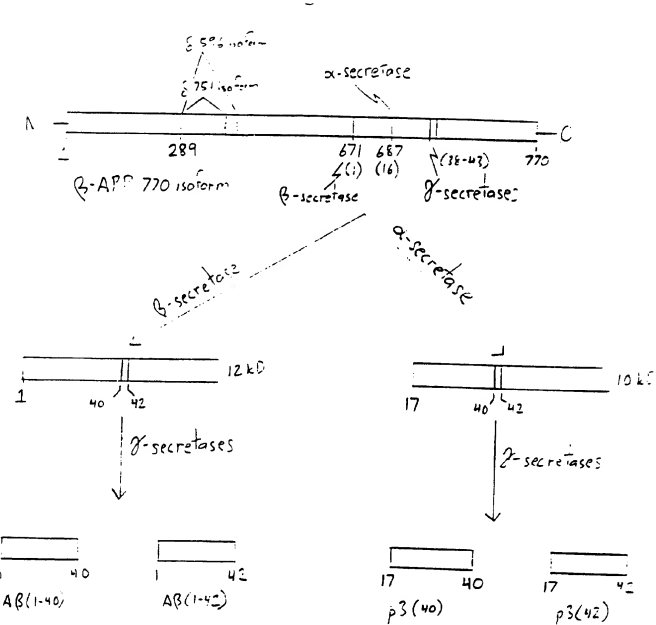
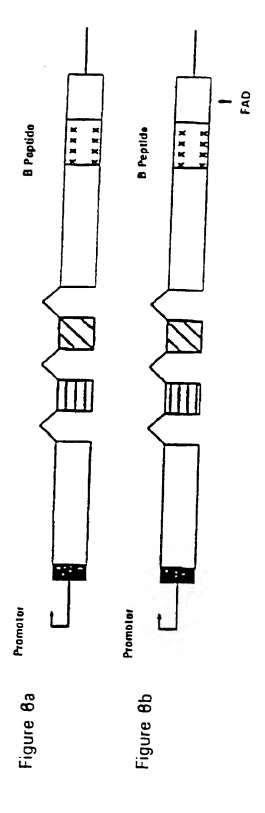


Fig 6

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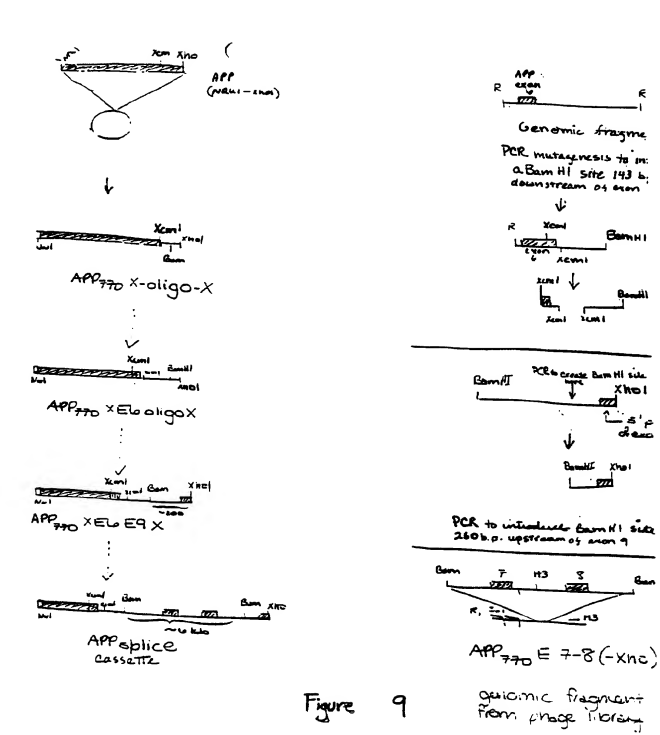


FIG 10A

(2)	(i;) (ii;) (iv) (ix)) SE () () () () () () () () () () () ()	QUENCA LIBO TO STATUTE TI - STATUTE ATURE	AME / I	HARA: NUC. NUC. DEDNI OGY: YPE: AL: NO KEY: INFO	CTER 310 leic ESS: lin cDN NO CDS 1-2: DRMA	ISTI base aci dou ear A	CS: pai: d ble :/fi	unct:	ion= D:5:	°co:	ding	reg	ion :	for .	APP77 0.	N
ATG Met 1	CTG Leu	CCC Pro	GGT Gly	TTG Leu 5	GCA Ala	CTG Leu	CTC Leu	CTG Leu	CTG Leu 10	GCC Ala	GCC Ala	TGG Trp	ACG Thr	GCT Ala 15	CGG Arg		48
GCG Ala	CTG Leu	GAG Glu	GTA Val 20	CCC	ACT Thr	GAT Asp	Gly	AAT Asn 25	GCT Ala	GŢĀ GGC	CTG Leu	CTG Leu	GCT Ala 30	GAA Glu	CCC		96
CAG Gln	ATT Ile	GCC Ala 35	ATG Met	TTC Phe	TGT Cys	GGC Gly	AGA Arg 40	CTG Leu	AAC Asn	ATG Met	CAC His	ATG Met 45	AAT Asn	GTC Val	CAG Gln		144
AAT Asn	GGG Gly 50	AAG Lys	TGG	GAT Asp	TCA Ser	GAT Asp 55	CCA Pro	TCA Ser	GGG Gly	ACC Thr	AAA Lys 60	ACC Thr	TGC Cys	ATT Ile	GAT Asp		192
ACC	AAG	GAA	GGC	ATC	CTG	CAG	TAT	TGC	CAA	GAA	GTC	TAC	CCT	GAA	CTG		240

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Th 6	r Ly 5	s Gl	u Gl	y Il	e Lei 7	u Gli	а Ту	т Су	s Gl	n Gl 7	u Va 5	l Ty	r Pr	o G1	u Leu	
CA Gl:	G AT n Il	C AC e Th	C AA	T GTO		A GAJ	A GCC	C AAG	T GT	A CC	_	G AC	C AT	C CA e Gl	BO G AAC n Asn	288
TG	G TG	C AA	G CG	3 660	- -	ስ አልር	- Cac	· •						9	5 T GTG e Val	336
ATT		TA	c cec	TGC	מדד י	بعدى							110	0		384
GTI	רכים	li: GAC	5 AAG	י דיקר	מממי		120	Cha	, ene	· vali	Ser	125	Ala	a Lei	u Leu	301
	130)	1-	-7-	-,-	135	Deu	nis	GID	. GIU	140	Met	Ası	Va.	i Cys	432
145					150	****	1111	VAI	ATA	Lys 155	Glu	Thr	Cys	Ser	GAG Glu 160	480
_				165	*****	qen	+ y L	GIY	170	Leu	Leu	Pro	Сув	G1y 175		528
•	-,-		180	J. y	V4.	GIU	Lite	185	Cys	Cys	Pro	Leu	Ala 190	Glu		576
		195	GTG Val	wah	361	ALE	200	ATE	GIU	Glu	Asp	Asp 205	Ser	Asp	Val	624
•	210		GGA Gly	~~ a	w.p	215	жар	Tyr	AIA	Asp	Gly 220	Ser	Glu	Asp	Lys	672
225			GTA Val	716	230	GIU	GIU	GIU	VAI	235	Glu	Val	Glu	Glu	Glu 240	720
			GAT Asp	245	310	veħ	vab	GIU	250	GIY	Asp	Glu	Val	Glu 255	Glu	768
			GAA Glu 260		.,.	G.u.	G14	265	ing	GIU	Arg	Thr	Thr 270	Ser	Ile	816
		275	ACC Thr	- ***	* ***		280	GIU.	Ser	VAI	Glu	Glu 285	Val	Val	Arg	864
	290	-,2	TCT Ser			295	J14 .	131T /	GIY .	PIO	300	Arg .	Ala	Met	Ile	912
TCC Ser 305	CGC Arg	TGG Trp	TAC Tyr	F 11.E	GAT (Asp \ 310	GTG ; Val :	ACT (Thr (GAA (Glu (ara :	AAG '	TGT (GCC (Ala :	CCA Pro	TTC Phe	TTT Phe 320	960

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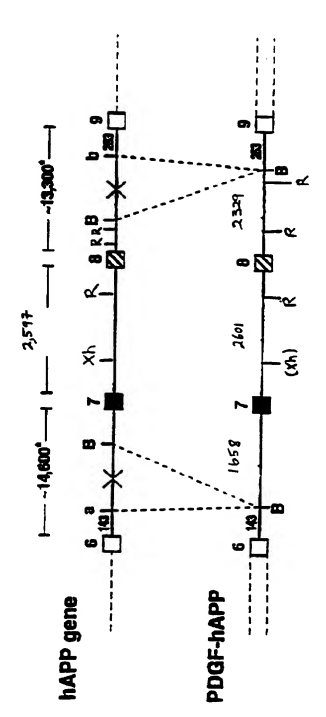
TAC	GG(GG) Gly	A TGT	GGC Gly 325	92	AAC Asn	CGC Arg	AA(Asi	0AA C 1 ABC 330	1 Phe	GA(C ACA P Thr	GAL Glu	GAG Gli Gli 335	TAC Tyr		1008
TGC	ATO Met	GCC Ala	Val	,-	GGC Gly	AGC Ser	GCC Ala	ATO Met	. ser	CAR Glr	AG7	TTA Leu	CTC Leu 350	Lys	ACT Thr		1056
		355		Dec	. Ala	. ALG	360	Pro) Val	Lys	Leu	Pro 365	Thr	Thr	GCA Ala		1104
GCC Ala	Ser 370		Pro	GAT Asp	GCC Ala	GTT Val 375	GAC Asp	AAG Lys	TAT Tyr	CTC Leu	GAG Glu 380	Thr	CCT	GGG Gly	GAT Asp		1152
GAG Glu 385	*****	GAA Glu	CAT His	GCC Ala	CAT His 390	TTC Phe	CAG Gln	AAA Lys	GCC Ala	AAA Lys 395	Glu	AGG Arg	CTT Leu	GAG Glu	GCC Ala 400		1200
J, S	uta	λig	GIU	405	Met	TCC Ser	GIN	Val	Met 410	Arg	Glu	Trp	Glu	Glu 415	Ala	:	1248
	7+9	9111	420	Lys	ASII	TTG Leu	PIO	125	Ala	Asp	Lys	Lys	Ala 430	Val	Ile	:	1296
G11	nis	435	GIN	GIU	LVS	GTG Val	440	Ser	Leu	Glu	Gln	Glu 445	Ala	Ala	Asn	1	1344
GIU	450	GIII	Gin	ren	Val	GAG Glu 455	Thr	HIS	Met	Ala	Arg 460	Val	Glu	Ala	Met	1	.392
465	wei!	veh	Arg	Arg	470	CTG Leu	AIA	Leu	Glu	Asn 475	Tyr	Ile	Thr	Ala	Leu 480	1	440
GIII	AIA	VAI	PIO	485	Arg	CCT Pro	Arg	HIS	Val 490	Phe	Asn	Met	Leu	Lys 495	Lys	1	488
- 7 -	val	AIG	500	GIU	GID	AAG Lys	ABD	Arg 505	GIN	His	Thr	Leu	Ly s 510	His	Phe	1	536
G. u	uiè	515	Arg	Mec	VAI	gat Asp	Pro 520	Lys	Lys	Ala	Ala	Gln 525	Ile	Arg	Ser	1	584
G 1 11	530	MEL	Inc	WIR	ren	CGT Arg 535	VAI	116	Tyr	Glu	Arg 540	Met	Asn	Gln	Ser	1	632
545	SET	Den	Leu	Tyr	550	GTG (Val)	Pro .	ALA	Val	A1a 555	Glu	Glu	Ile	Gln	As p 560	1	6 B O
G AA Glu	GTT Val	GAT Asp	GAG Glu	CTG Leu	CTT Leu	CAG A	AAA Lys	GAG Glu	CAA Gln	AAC Asn	TAT Tyr	TCA Ser	GAT Asp	GAC Asp	GTC Val	1	728

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				565					570	_				575		
Leu	ALA	Asn	580	Ile	Ser	Glu	Pro	Arg 585	Ile	Ser	Tyr	Gly	Asn 590	Asp	Ala	1776
CTC Leu	ATG Met	CCA Pro 595	TCT Ser	TTG Leu	ACC	GAA Glu	ACG Thr 600	AAA Lys	ACC	ACC Thr	GTG Val	GAG Glu 605	CTC Leu	CTT Leu	CCC	1824
GTG Val	AAT Asn 610	GGA Gly	GAG Glu	TTC Phe	AGC Ser	CTG Leu 615	GAC Asp	GAT Asp	CTC	CAG Gln	CCG Pro 620	TGG Trp	CAT His	TCT Ser	TTT Phe	1872
GGG Gly 625	GCT Ala	GAC Asp	TCT Ser	GTG Val	CCA Pro 630	GCC Ala	AAC Asn	ACA Thr	GAA Glu	AAC Asn 635	GAA Glu	GTT Val	GAG Glu	CCT Pro	GTT Val 640	1920
GAT Asp	GCC Ala	CGC Arg	CCT	GCT Ala 645	GCC Ala	GAC Asp	CGA Arg	GGA Gly	CTG Leu 650	ACC Thr	ACT Thr	CGA Arg	CCA Pro	GGT Gly 655	TCT Ser	1968
GGG Gly	TTG Leu	ACA Thr	AAT Asn 660	ATC Ile	AAG Lys	ACG Thr	GAG Glu	GAG Glu 665	ATC Ile	TCT Ser	GAA Glu	GTG Val	AAG Lys 670	ATG Met	GAT Asp	2016
GCA Ala	GAA Glu	TTC Phe 675	CGA Arg	CAT His	GAC Asp	TCA Ser	GGA Gly 680	TAT Tyr	GAA Glu	GTT Val	CAT His	CAT His 685	CAA Gln	AAA Lys	TTG Leu	2064
GTG Val	TTC Phe 690	TTT Phe	GCA Ala	GAA Glu	GAT Asp	GTG Val 695	GGT Gly	TCA Ser	AAC Asn	AAA Lys	GGT Gly 700	GCA Ala	ATC Ile	ATT Ile	GGA Gly	2112
CTC Leu 705	ATG Met	GTG Val	GGC Gly	GGT Gly	GTT Val 710	GTC Val	ATA Ile	GCG Ala	ACA Thr	GTG Val 715	ATC Ile	GTC Val	ATC Ile	ACC Thr	TTG Leu 720	2160
GTG Val	ATG Met	CTG Leu	AAG Lys	AAG Lys 725	AAA Lys	CAG Gln	TAC Tyr	ACA Thr	TCC Ser 730	ATT Ile	CAT His	CAT His	GGT Gly	GTG Val 735	GTG Val	2208
GAG Glu	GTT Val	GAC Asp	GCC Ala 740	GCT Ala	GTC Val	ACC Thr	CCA Pro	GAG Glu 745	GAG Glu	CGC	CAC His	CTG Leu	TCC Ser 750	aag Lys	ATG Met	2256
GIN	Gln	AAC Asn 755	GGC Gly	TAC Tyr	GAA Glu	Asn	CCA Pro 760	ACC Thr	TAC Tyr	AAG Lys	TTC Phe	TTT Phe 765	GAG Glu	CAG Gln	ATG Met	2304
CAG Gln																2310

Note: distances are not done to scate

Numbers indicate distances in base pairs

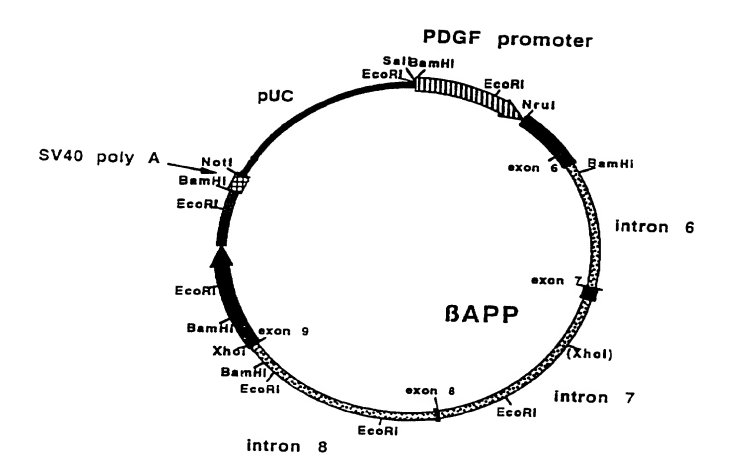


a, b - sites of mutagenesis where Bemtl sites were engineered for deletion B = location of Bam HI sites
X = xho! (x) Xho! site

R= 800 R1

intron 7 was expanded by 4 bp. when the xhol site was distroyed

Figure 12



PDAPP vector map

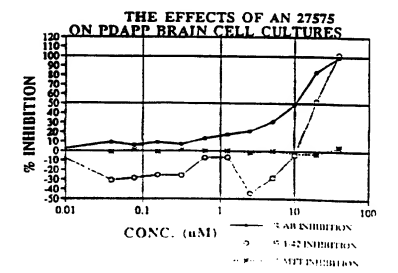
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FIG. 13A

S MITT INHIBITION

4 MTT INTIBITION



THESE POAPP CULTURES WERE GROWING LARSS

FIG. 13B

FIG. 13C

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0.01

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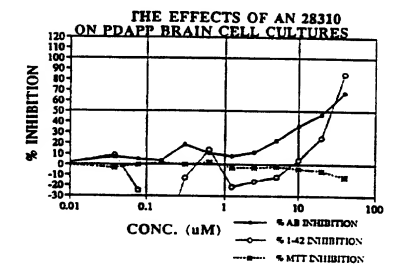
120: 110: 100: 90: 80: 70: 60: 50: 40: 30: 20: 10: % INHIBITION 0 -10

FIG. 13D

0.1 10 100 S AB INTERITION CONC. (uM) 4 1-42 INHIBITION

S MIT DIHERTION

FIG. 13E



120-110-100-90-80-70-60-50-40-20-10-% INHIBITION -10--20--30-0.01 0.1 10 100 % AB INHIBITION CONC. (uM) ····O···· % 1-42 INHIBITION

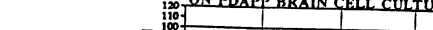
THE EFFECTS OF AN 28313

FIG. 13F

^{*} THESE PDAPP CULTURES WERE GROWN IN 9% FBS

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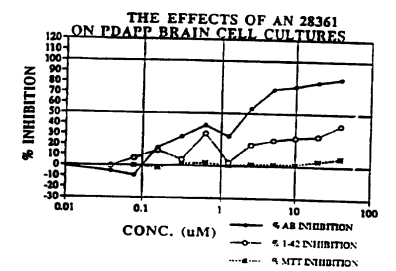
THE EFFECTS OF AN 28358 ON PDAPP BRAIN CELL CULTURES 120-110-100-90-80-70-60-50-10--10--20--30-% INHIBITION 0.01 0.1 10 100 NOTIFIELD EA P CONC. (uM)

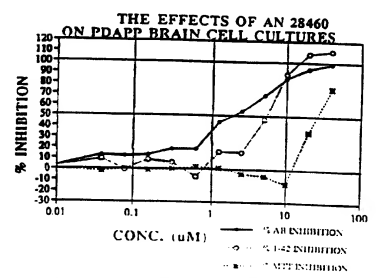
€ 1-42 EXTERITION S MIT INHERITION

FIG. 13G

FIG. 13H

FIG. 131





THESE PDAPP CULTURES WERE GROWTHIN 9% FBS



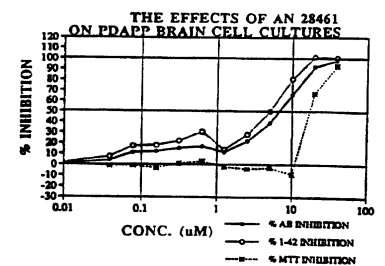


FIG. 13K

FIG. 13J

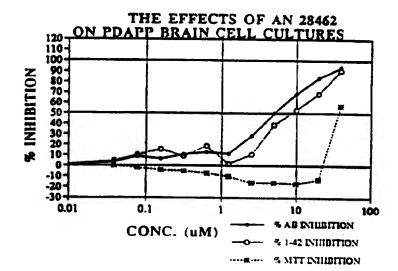
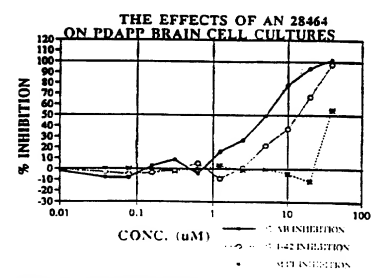


FIG. 13L



^{*} THESE PDAPP CULTURES WERE GROWNING CARS

FIG. 13M

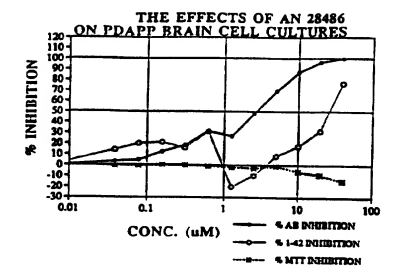


FIG. 13N

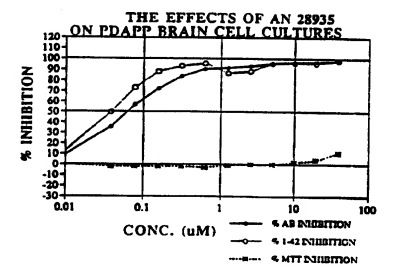
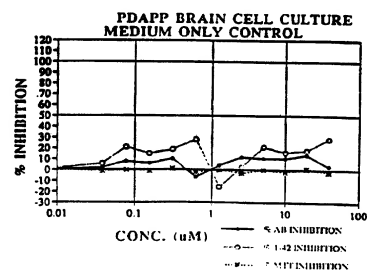


FIG. 130



THESE PDAPP CULTURES WERE GROWN IN 9% FBS

INTERNATIONAL SEARCH REPORT

International Application No PCT/US 97/10601

			,
A. CLASS IPC 6	G01N33/68 G01N33/50		
According t	to International Patent Classification (IPC) or to both national class	sification and IPC	
B. FIELDS	SEARCHED		****
Minimum d IPC 6	ocumentation searched (classification system followed by classific $GO1N$	cation symbols)	
Documenta	ation searched other than minimum documentation to the extent th	at such documents are included in the fields se	earched
Electronic o	data base consulted during the international search (name of data	base and, where practical, search terms used	3)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
X	WO 91 04339 A (CALIFORNIA BIOTE INCORPORATED) 4 April 1991 see claims 1,10-14; example 5	CHNOLOGY	1-4
A	WO 94 01772 A (THE CHILDREN'S M CENTER CORPORATION) 20 January see claim 1		1-58
Α	WO 93 16101 A (ATHENA NEUROSCIE INCORPORATED & ELI LILLY AND CO August 1993 see claim 1		1-58
А	WO 94 10569 A (D.B. SCHENK ET A 1994 see claim 1 	L.) 11 May	1-58
Furti	her documents are listed in the continuation of box C.	χ Patent family members are listed	in annex.
"A" docume consider of filing docume which citation "O" docume other r	ont which may throw doubts on priority claim(s) or is cited to establish the publication date of another in or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or	"T" later document published after the inte or priority date and not in conflict with cited to understand the principle or the invention "X" document of particular relevance; the cannot be considered novel or cannot lovolve an inventive step when the do "Y" document of particular relevance; the cannot be considered to involve an indocument is combined with one or ments, such combination being obvious the art. "&" document member of the same patent	the application but secry underlying the claimed invention it be considered to countent is taken alone claimed invention iventive step when the ore other such docu-us to a person skilled
Date of the	actual completion of theinternational search	Date of mailing of the international sea	
	9 November 1997	27/11/1997	
Name and n	nailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,	Authorized officer	

INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No
PET/US 97/10601

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9104339 A	04-04-91	AU 641434 B AU 6431190 A CA 2065404 A EP 0493470 A JP 5502368 T US 5221607 A	23-09-93 18-04-91 19-03-91 08-07-92 28-04-93 22-06-93
WO 9401772 A	20-01-94	NONE	
WO 9316101 A	19-08-93	US 5538845 A AU 3585293 A	23-07-96 03-09-93
WO 9410569 A	11-05-94	AU 4844493 A CA 2105903 A EP 0667959 A JP 8502587 T US 5593846 A	24-05-94 27-04-94 23-08-95 19-03-96 14-01-97